COMPUTER AIDED PERFORMANCE STUDY & COST ANALYSIS OF A DOUBLE-STAGE-VAPOUR ABSORPTION SYSTEM

Ву

ARUN MEHTA

ME 1987

M

MEH

Com



DEPARTMENT OF MECHANICAL ENGINEERING

INDIAN INSTITUTE OF TECHNOLOGY, KANPUR

AUGUST, 1987

COMPUTER AIDED PERFORMANCE STUDY & COST ANALYSIS OF A DOUBLE-STAGE-VAPOUR ABSORPTION SYSTEM

A Thesis Submitted
In Partial Fulfilment of the Requirements
for the Degree of

MASTER OF TECHNOLOGY

By

ARUN MEHTA

to the

DEPARTMENT OF MECHANICAL ENGINEERING

INDIAN INSTITUTE OF TECHNOLOGY, KANPUR

AUGUST, 1987

CHE- 1907-M- MEH-COM

21 SEP 1987
CENTRAL LIBRARY
Acc. No. A 97902



CERTIFICATE

This is to certify that this work on Computer Aided Performance Study and Cost Analysis of a Double-Stage-Vapour-Absorption-System has been carried out under my supervision and it has not been submitted elsewhere for a degree.

H.C. AGRAWAL)
Professor

Department of Mechanical Engineering, Indian Institute of Technology, Kanpur

July, 1987

<u>ACKNOWLEDGEMENT</u>

With immense pleasure, I express my deep sense of gratitude and regards to Professor H.C. Agrawal, for inspiring me to take up this problem and providing his inspiring guidance and constructive criticism to overcome all difficulties encountered during this period. His intense interest and unfailing assistance have been the main factors responsible for the completion of this work in time.

I am very grateful to the members of the faculty at the Department of Mechanical Engineering for their valuable contribution in my education.

The work would have been very difficult without the extremely helpful and valuable assistance given by Mr. N.K. Patnaik, my colleague and friend. I am highly grateful and thankful to him. I wish to express my thanks, also to Mr. Amar J. Borzthakur who made my stay at this place unforgettable.

I am deeply beholden to the members of my family for their abiding inspiration and loving patronage.

The get up of the thesis reflects sincere and pains -- ta king efforts of several peoples. I am thankful in particular to Mr. H.V.C. Srivastava for excellent typing.

Arun Mehta

CONTENTS

			Page
CERTIFICA	ATE		i
ACKNOWLE	DGEMENT		ii
LIST OF	FIGURES		vi
LIST OF	PABLES		vii
NOMENCLA	TURE		ix
ABSTRACT		*	хi
I	V	VAPOUR ABSORPTION AND COMPRESSION	1
		SYSTEMS	
	1.1	INTRODUCTION	1
	1.2	LITERATURE -SURVEY	8
÷	1.3	PRESENT WORK	11
II	SINGLE	-STAGE-VAPOUR-ABSORPTION-SYSTEM	16
	2.1	DESCRIPTION OF THE SYSTEM	16
	2.2	THERMODYNAMIC ANALYSIS OF THE SYSTEM	17
	2.3	COMPUTER PROGRAMME PACKAGE	26
- III	A NIA T VC	IS AND COMPUTER MODELLING OF DOUBLE-	28
ماد ماد ماد د د د د د د د د د د د د د د د د د د د		VAPOUR-ABSORPTION-SYSTEM	2.0
**	3.1	DESCRIPTION OF THE SYSTEM	28
- 10 - 10 - 10 - 10 - 10 - 10 - 10 - 10	3.2	THERMODYNAMIC ANALYSIS OF THE SYSTEM	30
	3.3	COMPUTER MODELLING	38

	3 . 3.1	Subroutine for Solving State Equations	39
·	3.3.2	Algorith m, Newton-Raphson Technique	40
IV	ECONOMI	CS OF VAPOUR-ABSORPTION AND VAPOUR-	42
	COMPRE	CSSION SYSTEMS	
V	RESULT	S AND DISCUSSION	, 54
REFERE	ENCES		84
APPENT	OTCES	·	87

LIST OF FIGURES

Fig. No.	Description	Page No.
1.1	Flow Diagram of a Vapour-Compression- System	3
1.2	One-Stage Absorption System with Heat Exchanger	Lį.
1.3	Double-Effect Absorption Unit (The Trane Company)	12
1.4	Schematic Diagram of Double-Stage- Vapour-Absorption-System	13
2.1	Heat-Operated Refrigeration Cycle	25
2.2	Flow-Chart for Single-Stage-Absorption- System	27
3.1	Flow-Chart for Computer Modelling of a Double-Stage-Vapour-Absorption-System	41
5.1	Variation of C.O.P. with Generator-I Temperature ($T_c=20^{\circ}\text{C}$, 25°C ; $T_a=20^{\circ}\text{C}$, $T_e=5$	65 °C)
5.2	Variation of C.O.P. with Generator-I Temperature ($T_c=30^{\circ}$ C, $T_a=25^{\circ}$ C, $T_e=5^{\circ}$ C)	66
5.3	Variation of C.O.P. with Generator-I Temperature ($T_c=30^{\circ}$ C, $T_a=30^{\circ}$ C, $T_e=5^{\circ}$ C)	67

5.4	Variation of C.O.P. with Generator-I Temperature $(T_c=35^{\circ}C, T_a=30^{\circ}C, T_a=8^{\circ}C)$	68
5.5	Variation of C.O.P. with condenser- Temperature $(T_g=90^{\circ}\text{C}, T_a=25^{\circ}\text{C}, T_e=8^{\circ}\text{C})$	71
5.6	Variation of C.O.P. with Condenser- Temperature (Tg=95°C,Ta=25°C,Te=8°C)	72
5 . 7	Variation of C.O.P. with Evaporator- Temperature (Double-Stage, $T_c=25^{\circ}\text{C}$, $T_a=20^{\circ}\text{C}$)	75
5 . 8	Variation of C.O.P. with Absorber- Temperature (Double-Stage, Te=5°C, 8°C; Tg=90°C, Tc=30°C)	77
5.9	Variation of Generator-II Temperature with Generator-I Temperature	79
5.10	Variation of Generators Loadswith Generator-I Temperature	80
5.11	Variation of C.O.P. with Heat Exchanger Effectiveness	81

LIST OF TABLES

TABLE NO.	DESCRIPTION	PAGE NO
1	Effect of Generator-I Temperature	61
	on operating parameters	
	$(T_c = 20 {}^{\circ}\text{C}, T_a = 20 {}^{\circ}\text{C}, T_e = 5 {}^{\circ}\text{C},$	
	$\eta_1 = \eta_2 = 0.90$)	
2	Effect of Generator-I Temperature	62
	on operating parameters	•
t	$(T_c = 25 {}^{\circ}\text{C}, T_a = 20 {}^{\circ}\text{C}, T_e = 5 {}^{\circ}\text{C},$	
	$\eta_1 = \eta_2 = 0.90$)	
3	Effect of Generator-I Temperature	65
	on operating parameters	
	$(T_c = 30 {}^{\circ}\text{C}, T_a = 25 {}^{\circ}\text{C}, T_e = 5 {}^{\circ}\text{C},$	
	$\eta_1 = \eta_2 = 0.90$)	
4	Effect of Generator-I Temperature	64
	on operating parameters	
	$(T_c = 35 {}^{\circ}\text{C}, T_a = 30 {}^{\circ}\text{C}, T_e = 8 {}^{\circ}\text{C},$	
	$\eta_1 = \eta_2 = 0.90$)	
5	Effect of Condenser-Temperature on	69
	C.O.P.	
	$(T_g = 90 {}^{\circ}\text{C}, T_a = 25 {}^{\circ}\text{C}, T_e = 8 {}^{\circ}\text{C},$	
	$\eta_1 = \eta_2 = 0.90$)	
6	Effect of Condenser-Temperature on	70
	C.O.P.	
	$(T_g = 95 {}^{\circ}\text{C}, T_a = 25 {}^{\circ}\text{C}, T_e = 8 {}^{\circ}\text{C},$	and the second s
	$\eta_1 = \eta_2 = 0.90$)	
		1

7 Effect of Evaporator-Temperature on 75 C.O.P. $(T_g = 85 \text{ °C}, T_c = 25 \text{ °C}, T_a = 20 \text{ °C},$ $\eta_1 = \eta_2 = 0.90)$

8 Effect of Evaporator-Temperature on 74 C.O.P. $(T_g = 95 \text{ }^{\circ}\text{C}, T_c = 25 \text{ }^{\circ}\text{C}, T_a = 20 \text{ }^{\circ}\text{C},$ $\eta_1 = \eta_2 = 0.90)$

9 Effect of Evaporator-Temperature on 7^{l_1} C.O.P. $(T_g = 90 \text{ °C}, T_c = 30 \text{ °C}, T_a = 25 \text{ °C},$ $\eta_1 = \eta_2 = 0.90)$

Effect of Absorber-Temperature on the C.O.P. $(T_g = 90 \text{ }^{\circ}\text{C}, T_c = 30 \text{ }^{\circ}\text{C}, T_e = 50 \text{ }^{\circ}\text{C},$ $\eta_1 = \eta_2 = 0.90)$

Effect of Absorber-Temperature on the C.O.P. $(T_g = 90 \text{ °C}, T_c = 30 \text{ °C}, T_e = 8 \text{ °C},$ $\eta_1 = \eta_2 = 0.90)$

Effect of Heat-Exchanger effectiveness on 78 C.O.P. $(T_g = 90 \text{ °C}, T_c = 25 \text{ °C}, T_a = 20 \text{ °C}, T_e = 5 \text{ °C}$

NOMENCLATURE

A	Surface area	$[m^2]$
	Specific heat	[kJ/kg. K]
c _P	Diameter of tube	[m]
f	Friction factor	
g	Acceleration due to gravity	$[m/s^2]$
Н	Enthalpy	[kJ/kg]
h	Heat transfer coefficient	$[W/m^2.K]$
K	Thermal conductivity	[W/m.K]
ṁ̀	Mass flow rate	[kg/s]
N	Number of tubes per pass	
P	Pressure	[k Pa]
Q	Volume flow rate	[m ³ /s]
q	Heat load	[kW]
R	Thermal resistance	$[m^2.K/W]$
t	Temperature	[°c]
t _f	Temperature	[°F]
TR	Refrigerant temperature	[°c]
TS	Solution temperature	[°C]
U	Overall heat transfer coefficient	$[W/m^2.K]$

	ν	Velocity	[m/s]	
	Х	Lithium bromide concentration /.	[kg/kg o	f sol.]
	Subscrip	<u>ts</u>		
	а	Referred to absorber		
	С	Referred to condenser condition		
	е	Referred to evaporator condition		
	ff	Fouling condition		
	gi	Referred to first stage generator	·	
	g2	Referred to second stage generator		
	i	Referred to inlet condition		
	m	Mean		
	0	Referred to outlet condition		
•	t	Tube side		
	W	Water side		
	1,2,3,,,	7 State points referred to Fig. (1.3)	for singl	e-stage
	1,2,3,	18 Stage points referred to Fig. (1.4) for	•
		double-stage		
	I,II	First and second heat exchanger		
	Greek Sy	mbols:		
	μ	Dynamic Viscosity	[Pa.s]	
	ε	Absolute roughness	[mm]	
	ρ	Density of fluid	$[kg/m^3]$	
	ν	Specific volume	$[m^3/kg]$	
6	η	Heat exchanger effectiveness		
	- pr;			

ABSTRACT

The escalating cost of energy and the fast depleting sources of conventional—energy have sounded an alarm to the world, urging us to be meticulous in using any form of energy, at our disposal. Using waste heat energy from various industrial processes and the use of non-conventional energy sources, such as: solar, geothermal, bio-gas, wind, tidal etc., are vitally important propositions in this context. A double-stage-vapour-absorption-system to produce refrigeration, attempts to fulfil the efficient use of the above energy resources.

Keeping the above fact in mind, the present work is devoted to the analysis of a double-stage-vapour-absorption-system. The complete performance-study of the system is carried out to determine the variation of C.O.P. (the index of performance of the system) with various operating parameters. The optimum values of these parameters are also evaluated for maximum C.O.P. This C.O.P. is compared with the C.O.P. of the corresponding single-stage-vapour-absorption-system.

The most conventional method of producing refrigeration is by using vapour-compression-system, which uses the valuable mechanical energy for its operation. Therefore, is the major

competitor to the proposed system, presented in the thesis. Hence, a comparative cost-analysis is also undertaken in the present study to evaluate the relative economics of operation for the two systems.

A digital computer DEC-1090 has been used to help in carrying out the analysis. Required computer programmes have been developed to perform this analysis, which are made very general to encompass a class of similar problems. Results have been discussed and the following conclusions drawn:

- 1. The C.O.P. obtained in a double stage-system is quite high (1.785 maximum) as compared to single stage (0.84 maximum). However, very high input source-temperature is required for system's operation.
- 2. This system has very high values of C.O.P., almost approaching the ideal value at high condenser temperatures. The practicability of using high condenser-temperatures obviates the necessity of using a cooling tower along with the condenser and an air cooled condenser may be used for the purpose. This helps in a reduction in total cost of the system.
- J. A double-stage-vapour-absorption-system is more economical than a vapour-compression-system in terms of the annual running cost as it does not require compressor for its operation, which consumes most of the energy required for the operation of a vapour-compression-system.

CHAPTER-I

VAPOUR ABSORPTION AND COMPRESSION SYSTEMS

1.1 INTRODUCTION:

Vapour compression system is invariably used in most of the refrigeration and air conditioning industries. The outline of such a system along with all its basic components are shown in Fig. 1.1. However, it uses the costly mechanical energy in the form of compressor work and the ever-rising costs of this energy due to continuous depleting energy resources have demanded that this valuable energy should be conserved, as much and as far as possible. There are broadly two ways which can assist in conserving the conventional energy:

- (i) using non-conventional energy sources like solar, tidal, bio-gas, geothermal and wind energy, economically and
- (ii) using waste energy to recover some heat out of it, which otherwise would be wasted. Waste heat in the form of air and hot water from industries such as steel, chemical, textile, foundaries etc. can be considered as practical examples.

Vapour absorption system, which can work using either of the above two alternativenergy sources or their

combination, is one of the very old methods to produce cold. Single-stage-vapour-absorption-system was first used in U.S.A. during the civil war after the supply of natural ice had been cut off from North. Many improvements have been incorporated in the system over the years. The skeleton diagram of such an improved version of single-stage-vapour-absorption system is shown in Fig. 1.2. Since, the thesis work is devoted to the vapour-absorption system, we describe below the various components of this system:

(i) Generator

It is basically a heating vessel, which receives heat from the heat source and transfers it to the solution flowing inside the tubes. The source of heat can be steam, solarheat, bio-gas, hot-water, hot-air etc.

(ii) Condenser

It is a shell and tube type heat exchanger located after the generator. Cooling water to cool refrigerant flows inside the tubes, whereas, refrigerant condenses in the shell.

(iii) Absorber

It is a heat-rejecting vessel. The strong absorbent solution in the absorber is sprayed to absorb the low pressure and temperature refrigerant vapour, coming out of the evaporator. Heat of absorption, which is generated due to exothermic

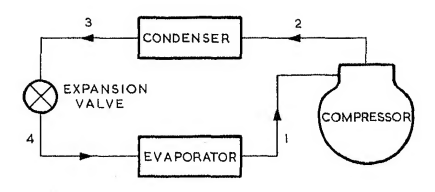


FIG. 1.1 FLOW DIAGRAM OF VAPOUR COMPRESSION SYSTEM.

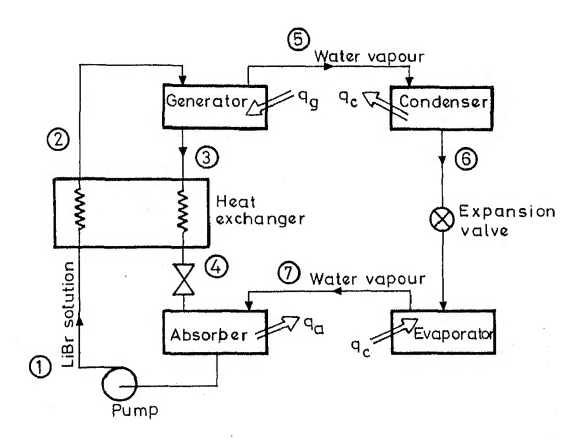


FIG. 1.2 ONE-STAGE ABSORPTION SYSTEM WITH HEAT EXCHANGER.

reaction, is rejected to the atmosphere, through the cooling water which flows inside the tubes.

(iv) Evaporator

Evaporator, sometimes also known as the Chiller, gives the desired cooling effect. Refrigerant flows inside the tubes, whereas liquid to be cooled flows over the tubes. Like condenser, it also has a shell and tube type construction.

(v) Solution-Heat-Exchanger

construction. It heats the weak solution on its path from the absorber to the generator by extracting heat from the relatively hotter strong solution returning from the generator. This reduces the amount of heat to be added to the solution in the generator.

(vi) Expansion-Device

Expansion-device i.e. expansion valve is used for the reduction of pressure and temperature of the refrigerant liquid, while flowing from the condenser to evaporator. The valve used is generally an orifice or other fixed restriction /11_7 type.

(vii) Solution Pump

It is generally electrical motor-driven centrifugalpump of hermatic design, which circulates the solution in the absorption system. The index of performance of a refrigeration system is represented by a dimensionless parameter termed as 'Coefficient of Performance', abbreviated as C.O.P.

For a vapour compression system, C.O.P. is defined as :

For a vapour absorption system, on the other hand, the C.O.P. is expressed as:

C.O.P. = Useful refrigerating effect Rate of heat addition in the generator

An absorption cycle has an appreciably lower value of C.O.P, than that of a compression cycle (0.8 versus 3.5).

A vapour compression cycle is work operated cycle (mechanical work in compressor), whereas, absorption cycle is heat operated cycle (heat addition in generator) and energy in the form of work is normally much more valuable and expensive than energy in the form of heat. It is, therefore, obvious that, the two systems cannot be measured on the same scale.

The vapour compression system pushed absorption system as to the charles are concerned.

A obsolete because of its being more reliable in operation and having better C.O.P. However, with the advent of worldwide concern over energy conservation, the vapour absorption system has again gained importance as it saves lot of

conventional, valuable, mechanical energy. This system has been found quite attractive in the space-conditioning applications, producing chilled water for various application and for preserving perishable commodities.

Two types of vapour absorption systems can be practised:

- (i) using Lithium Bromide as the absorbent and water as the refrigerant and
- (ii) using water as the absorbent and Ammonia as the refrigerant.

The combination adopted in the second type is called aqua-ammonia and was used in absorption systems, years before the LiBr-H₂O combination became popular, which has now almost completely replaced the aqua-ammonia system due to some of its following distinct advantages:

- (i) this combination yields higher C.O.P. at most operating temperatures.
- (ii) it operates at lower system pressures, thereby, requiring low pumping power.
- (iii) it can be used in commercial and residential buildings without any restriction from safety consideration.
- (iv) lower first cost due to the absence of parts like rectifier and analyzer.

Various improvements and refinements in conventional old Single-stage-absorption-system have been introduced from time to time to make vapour absorption system more efficient. It includes the reducing the generator temperature for system operation, increasing the allowable cooling water temperature and increasing the coefficient of performance, which, in turn, improves the efficiency of the system and conserves the energy.

The practicability of increasing the cooling water temperature is of special importance, because a sufficient reduction in the cost of the system may be achieved by using an air-cooled heat exchanger for heat rejection from the sinks of the absorption system and the requirement of a cooling tower may be eliminated.

1.2 LITERATURE-SURVEY

According to available literature tests for proving feasibility of solar air conditioning appeared in 1960 /1_7. Design of one ton solar operated LiBr-H₂O air conditioning system was presented at National Solar Energy Convection at Bhavnagar /2_7 and performance prediction of LiBr-H₂O absorption air conditioning system, utilizing the solar energy was put forward at International Solar Energy Congress at New Delhi /3_7.

Design and optimization of an absorption cycle with LiBr-H₂O and H₂O-NH₃ combination was studied by Alizadeh.et.al./4_7. This study shows that, in general, for fixed initial conditions and given system-refrigeration-capacity, higher generator temperature causes higher cooling ratio with smaller heat exchange surfaces and, consequently, lower cost. A comparison of the two cycles also indicates that LiBr-H₂O system is much simpler than NH₃-H₂O system and operates at a higher cooling ratio and for smaller heat exchange surfaces for the same conditions.

An economic evaluation of the use of Bio-gas to run a LiBr-H₂O absorption system has been carried out by Siddiqui et.al. <u>/</u>6_7. Coefficient of Performance of the system has been studied for various values of generator-temperature, pressure, evaporator-temperature, precooler and preheater effectiveness.

A dynamic model of solar air conditioning, including the solar collector and cooling tower, in terms of design parameters, is described by S.L. Grassie __7_7. It has the special feature that it provides a refrigerant storage tank associated with the generator to stabilize the flow.

absorption-system. Due to escalating energy cost, in recent times, more attention, however, is being paid towards the energy conservation and its effective utilization. This has necessitated the study of a more efficient and improved system. Double-stage-vapour-absorption-system is such an improvement, which is supposed to work on higher C.O.P. than a single stage with certain other advantages like reduced condenser load, material saving in the tubes etc.

one of the recent works on the study of a double-stage-vapour-absorption-system is done by Kaushik and Kumar in the year 1985 [8.7], in which they have considered the refrigerant-absorbent combination of NH3-H2O and NH3-LiNO3. The system consists of coupling of two conventional absorption cycles, so that the first stage evaporator produces cooling water to be circulated in the absorber of the second stage. The effect of operating variables, such as generator-temperature, evaporator-temperature and condenser-temperature on the C.O.P., heat transfer rates and relative circulation rates, has been

studied. Reference has been made by Stoecker and Jones <u>[9_7]</u> to the design of a double-stage-vapour-absorption-system proposed by the Trane Company of U.S.A. LiBr-H₂O combination has been used as the working fluid for the system. A circuit diagram is presented (Fig. 1.3). C.O.P., of the order of 1.0 to 1.4 has been reported for such a system.

1.3 PRESENT WORK

The present work is an extension of the design concept for the double-stage-vapour-absorption-system, enunciated by the Trane Company <u>19_7</u>. The same working fluid i.e. LiBr-H₂O combination is used. An improved design of the system has been proposed. The detailed circuit diagram for the same is shown in (Fig. 1.4). This design has the following merits in comparison with the one, given by the Trane Company.

the steam from generator-I, which drives off some water vapour from the LiBr-H₂O solution. Hence, the solution leaving generator-II, is a two-phase mixture. This two-phase mixture is then throttled to the condenser pressure in throttling-valve-I. Since, the throttling valves are not designed to handle two phase mixture, this operation is inefficient. On the other hand, in the improved system, presented in Fig.(1.4), no throttling of LiBr-H₂O solution is necessary, as generator-II is already maintained at condenser pressure.

(ii) In the system, developed by the Trane Company, generators-I and II are put in the same vessel. The thermodynamic properties of the LiBr-H₂O solution are such that, they impose a restriction on the heat exchanger-II. The level of cooling to be achieved in this heat exchanger for the LiBr-H₂O solution, entering the generator-II should be sufficient to accept heat from the steam, given off from generator-I. But in the system, proposed in the present work Fig. 1.4), generator-II is uncoupled from generator-I and, therefore, no restriction on the cooling capacity of the heat exchanger need be imposed.

The performance characteristics of the improved system are studied by the use of numerical modelling of the system which is based on the corresponding mass, energy and material balance equations of each component of the system. Effect of various component's temperature viz generator, absorber, condenser and evaporator on the performance of the system has been studied. Optimization of generator-I temperature for maximum C.O.P. along with minium generator-temperature required to carry out system operation have been determined. Effect of heat-exchanger effectiveness on the system performance has also been studied.

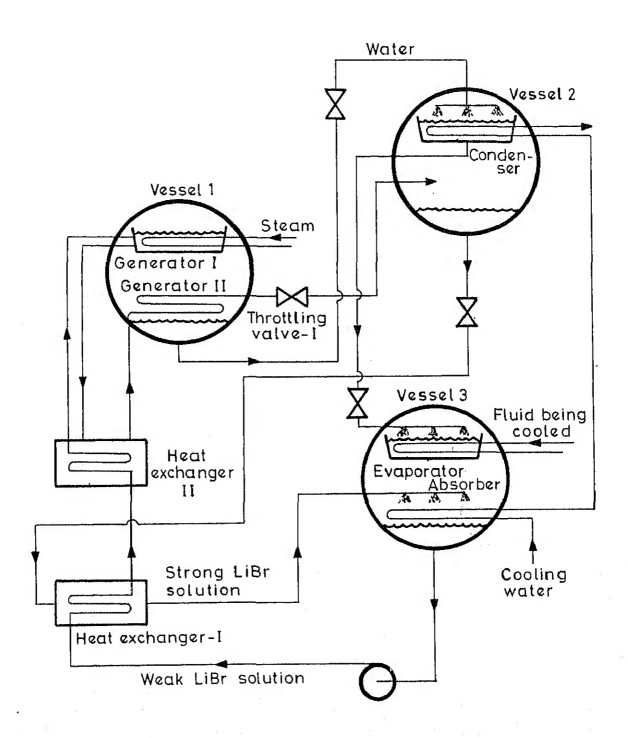


FIG. 1.3 DOUBLE-EFFECT ABSORPTION UNIT (THE TRANE COMPANY).

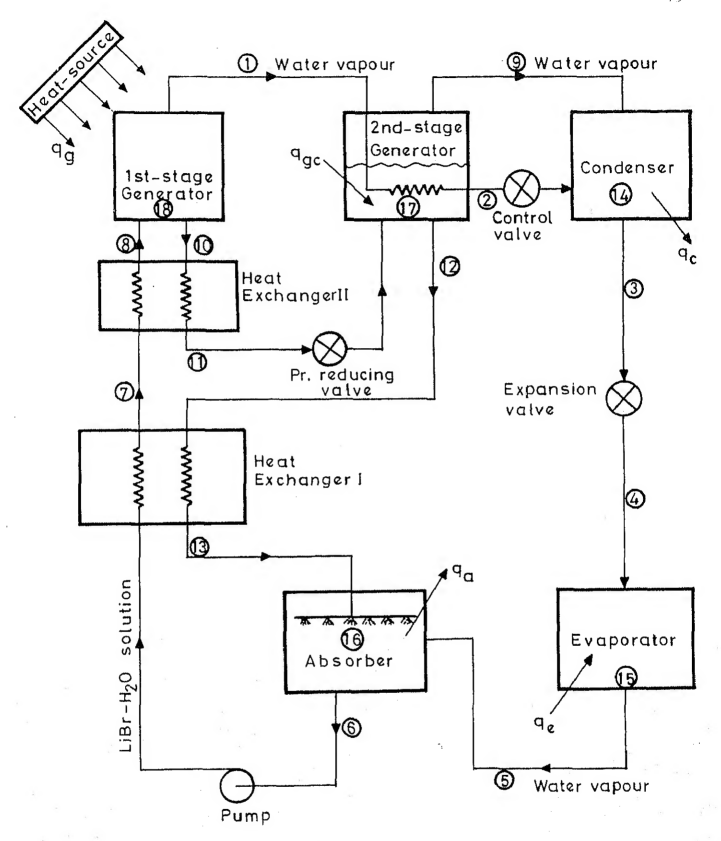


FIG. 1.4 SCHEMATIC DIAGRAM OF A DOUBLE-STAGE VAPOUR ABSORPTION SYSTEM (PRESENT WORK).

Comparison of the performance of this system is carried out with that of corresponding single-stage-vapour-absorption-system.

General computer programmes in FORTRAN language have been developed for both the single-stage and double-stage systems to evaluate the operational parameters (pressure, temperature, concentration, heat flow rates, mass flow rates of solution, refrigerant etc.), where-ever needed.

To emphasize the economic advantages of the double-stage LiBr-H₂O vapour absorption system, it's yearly operating cost has been compared with that of the vapour-compression-system of the same capacity. Suitable computer programme has been developed for this purpose. Operating cost includes the cost of power used in pumping the LiBr-H₂O solution and cooling water and also in raising the pressure level of the LiBr-H₂O solution from the absorber to the generator-I (Fig. 1.4).

For drawing graphs, 'LOTUS' subroutine in IBM personal computer has been used.

CHAPTER-II

SINGLE-STAGE-VAPOUR-ABSORPTION SYSTEM

2.1 DESCRIPTION OF THE SYSTEM:

The basic cycle of single-stage LiBr-H₂O vapour-absorption-system (Fig.1.2) consists of four temperature levels and two pressure level-zones.

Temperature level zones are :

- (i) Generator (T_g)
- (ii) Condenser (T_C)
- (iii) Absorber (T_a)
 - (iv) Evaporator (T_e)

Pressure level zones are :

- (i) the low pressure prevailing in the evaporator and absorber.
- (ii) the high pressure prevailing in the generator and condenser.

In the system operation, the weak solution of LiBr-H₂O is pumped from the absorber to generator through a solution heat exchanger. In the generator, the solution is heated at high temperature (60-90°C), so that, some of the refrigerant i.e. water is vaporised from the LiBr-H₂O solution. The remaining solution, which is strong solution (due to lower percentage of water) returns back to the absorber through

the heat exchanger. In the heat exchanger, this solution transfers some of its heat to the weak solution, pumped from the absorber. A throttling valve is also provided to develop a pressure drop between generator and absorber $(p_g > p_a)$.

The refrigerant vapour leaving the generator, passes to the condenser and is condensed, rejecting the heat of condensation to the circulating cooling water. The refrigerant-liquid in the condenser passes through the expansion valve and evaporates in the evaporator at very low pressure and temperature, extracting latent heat of vaporisation from the surroundings. It causes the cooling effect in the surrounding space. This vapour is then dissolved in the strong solution, returning back from the generator to absorber. This process produces heat of solution, which is carried away by the cooling water. After absorption, the weak solution is again pumped back to the generator. This completes the cycle of operation.

2.2 THERMODYNAMIC ANALYSIS OF THE SYSTEM :

A detailed thermodynamic analysis of the single-stageabsorption cycle has been carried out to evaluate the C.O.P. of the system. The following assumptions have been made:

- (i) the system is in a steady-state.
- (ii) the properties of the solution leaving a component are representative of the corresponding solutionproperties in the component.

- (iii) pressure loss in the heat exchanger is neglected.
- (iv) minimum terminal temperature difference is 5° C in the heat exchanging components of the system.
- (v) pressures in the evaporator and condenser are equal to the vapour pressure of the refrigerant i.e. water, at the corresponding evaporator and condenser temperatures.

Thermodynamic Analysis:

- (1) The absorber-temperature and the condenser-temperature are fixed by the available cooling water temperature. The requirements of the conditioned space fixes the evaporator-temperature. Heat exchanger effectiveness is assumed to be 0.9 which is in the commonly used range of its value \(\frac{10}{0} \). The available heat-source-temperature determines the generator-temperature.
- (2) The generator and the condenser, being directly connected to each other, exhibit the same pressure in them, i.e. $p_g = p_c$. For similar reason, pressures in the evaporator and the absorber are also equal i.e. $p_e = p_a$.

The condenser pressure 'p' as a function of condenser temperature 'T' is given in expression (A-2) of Appendix-1 which describes the state equations

for aqueous LiBr solution. Since ${}^tT_c^i$ is already known, ${}^tP_c^i$ can be evaluated using (A-2). Similarly, from the prior knowledge of ${}^tT_e^i$, ${}^tP_e^i$ can be determined.

- Using the already known values of absorber pressure and temperature, the concentration of LiBr-H₂O solution, leaving the absorber ${}^{1}X_{1}^{1}$ can be read from the graphs given in Appendix-2. Similarly, ${}^{1}X_{3}^{1}$, the concentration of the LiBr-H₂O solution, leaving the generator is obtained from the knowledge of generator pressure and temperature. Further as there is no change of concentration in the heat exchanger, we must have $X_{1}=X_{2}$ and $X_{3}=X_{4}$.
- (4) The conservation of mass for the working fluid in the generator is given by ;

rate of inflow of total mass to the generator (\bar{m}_2) = rate of outflow of total mass from the generator $(\bar{m}_3 + \bar{m}_5)$

or,

$$\dot{m}_2 = \dot{m}_3 + \dot{m}_5 \tag{2.1}$$

Similarly, conservation of LiBr in the generator requires,

$$\dot{m}_2 X_2 = \dot{m}_3 X_3$$
 (2.2)

At this stage, a term, called the Absorption-Recirculation-Ratio (ARR) is defined. 'ARR' is the mass of solution, pumped to the generator per unit mass of water vapour, supplied to the condenser. The 'ARR' is represented in terms of mathematical expression as:

$$ARR = \frac{\mathring{m}_2}{\mathring{m}_5} \tag{2.3}$$

Using (2.1) and (2.2), the expression for ARR reduces to,

ARR =
$$\frac{X_3}{X_3 - X_2}$$
 (2.4)

(5) The conservation of total mass in the absorber gives,

$$\dot{m}_1 = \dot{m}_4 + \dot{m}_7$$
 (2.5)

Also, the conservation of LiBr in the absorber yields,

$$\dot{m}_1 X_1 = \dot{m}_4 X_4$$
 (2.6)

Another term, called the Generator-Recirculation-Ratio (GRR) is defined. 'GRR' is the ratio of mass of LiBr-H₂O solution, leaving the generator to the mass of water-vapour driven off from the generator.

Mathematically, 'GRR' is given by,

$$GRR = \frac{\dot{m}_3}{\dot{m}_5} = \frac{\dot{m}_4}{\dot{m}_7}$$
 (2.7)

Using relations (2.5) and (2.6), the expression (2.7) reduces to ,

GRR =
$$\frac{X_1}{X_4 - X_1}$$
 (2.8)

The enthalpies of the fluid at various state points are to be determined in order to determine the heat loads on the various components in the system. In this context, enthalpies for LiBr-H₂O solution are obtained from the graph in Appendix-3 and the steam table <u>[9]</u>7 is used to get the enthalpy values for the refrigerant. Adopting this method, enthalpies at state points '1', '3', '5', '6' and '7' (Fig. 1.2) can be directly obtained. The corresponding enthalpies at state points '4', '2' are determined as following using the enthalpy values obtained above:

(a) Determination of H4:

Heat-exchanger effectiveness is defined as :

n = Enthalpy drop in the heat exchanger
Maximum possible enthalpy drop in the same

or,

$$n = \frac{H_3 - H_4}{H_3 - H_1} \tag{2.9}$$

In the above expression, all the parameters are known, except $H_{L\!\!\!/}$. Hence, $H_{L\!\!\!/}$ can be evaluated by the relation :

$$H_4 = H_3 - n (H_3 - H_1)$$
 (2.10)

(b) Determination of H2:

Energy-balance in the heat exchanger gives,

$$\dot{m}_3(H_3-H_4) = \dot{m}_1 (H_1-H_2)$$
 (2.11)

Since \dot{m}_3 and \dot{m}_1 are not known exclusively, both sides of expression (2.11) are divided by \dot{m}_5 . With this modification, expression (2.11) reduces to ,

$$\frac{\dot{m}_3}{\dot{m}_5} (H_3 - H_4) = \frac{\dot{m}_1}{\dot{m}_5} (H_1 - H_2)$$
 (2.12)

Using expression (2.3) and (2.7) in (2.12), we get,

$$GRR (H_3-H_4) = ARR (H_2-H_1)$$

or,

$$H_2 = \frac{GRR}{ARR} (H_3 - H_4) + H_1$$
 (2.13)

- (7) The heat transfer rates to and from the various system components can now be calculated by performing energy balance for each component:
 - (a) The heat flow rate in the evaporator:

$$q_e = H_7 - H_6$$
 (2.14)

(b) Heat, to be rejected in the condenser:

$$q_c = H_5 - H_6$$
 (2.15)

(c) Heat, to be supplied in the generator :

$$q_g = \dot{m}_5 H_5 + \dot{m}_3 H_3 - \dot{m}_2 H_2$$
 (2.16)

Assuming \dot{m}_5 as unity and dividing both sides of the expression (2.16) by \dot{m}_5 , we obtain,

$$q_g = H_5 + GRR \times H_3 - ARR \times H_2$$
 (2.17)

(d) Heat, to be rejected by the absorber :

$$q_3 = \mathring{m}_4 H_4 + \mathring{m}_7 H_7 - \mathring{m}_1 H_1$$
 (2.18)

Again, with necessary simplifications, we get,

$$q_{a} = GRR \ X \ H_{A} + H_{7} - ARR \ X \ H_{1}$$
 (2.19)

(8) For the total balance of energy in the system,

Total heat addition = Total heat rejection
or,

$$q_e + q_e = q_c + q_a$$
 (2.20)

This condition is very much essential, to be satisfied.

The Coefficient of Performance of the system is

then given by

C.O.P. =
$$\frac{q_e}{q_g}$$
 (2.21)

The value of C.O.P., obtained by the above procedure is then compared with the C.O.P. of the ideal system to get an idea of how closely the system approaches the ideal cycle. The procedure to obtain C.O.P. of the ideal system is given below:

An ideal vapour-absorption-cycle can be considered as a combination of a power cycle and refrigeration cycle (Fig. 2.1). The power cycle receives energy in the form of heat $\mathbf{q}_{\mathbf{g}}$ at an absolute temperature $\mathbf{T}_{\mathbf{g}}$, delivers some energy W in the form of work to the refrigeration cycle and rejects a quantity of energy $\mathbf{q}_{\mathbf{a}}$ in the form of heat at a temperature $\mathbf{T}_{\mathbf{a}}$. The refrigeration cycle receives the work W which is pumped along with the heat $\mathbf{q}_{\mathbf{e}}$ at the refrigerating temperature of $\mathbf{T}_{\mathbf{e}}$ to a temperature $\mathbf{T}_{\mathbf{c}}$, where the quantity $\mathbf{q}_{\mathbf{c}}$ is rejected.

The power-cycle, shown in Fig. (2.1), operates between the higher temperature limit T_g and the lower temperature limit T_a . Since, the analysis is carried out for an ideal cycle, the ideal thermal efficiency of this cycle is given by the Carnot's efficiency, which is represented as:

$$n_{\text{Power-cycle}} = \frac{W}{q_g}$$

$$= \frac{T_g - T_a}{T_g} \qquad (2.22)$$

Similarly, the refrigeration-cycle (Fig. 2.1), operates between sink temperature ${}^{'}T_{e}^{'}$ and source temperature ${}^{'}T_{c}^{'}$. For an ideal cycle, the C.O.P. is given by Carnot's C.O.P., represented as:

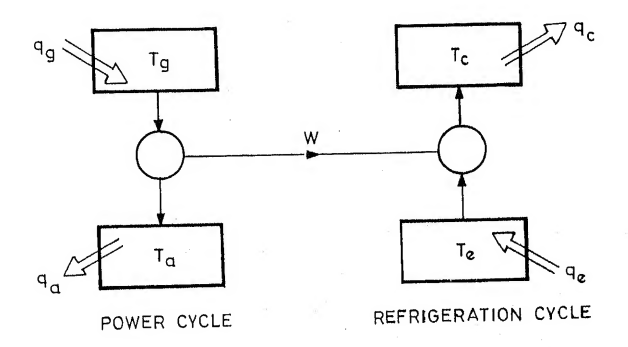


FIG. 2.1 HEAT-OPERATED REFRIGERATION CYCLE AS A COMBINATION OF A POWER CYCLE AND REFRIGERATION CYCLE.

$$C.O.P._{Ref} = \frac{q_e}{W}$$

$$= \frac{T_e}{T_c - T_e}$$
(2.23)

The C.O.P. of the entire cycle is the multiple of $n_{\mbox{Power-cycle}}$ and C.O.P. and is given by

C.O.P.Ideal =
$$n_{Power-cycle} \times C.O.P._{Ref}$$

= $\frac{W}{q_g} \times \frac{q_e}{W}$
= $\frac{q_e}{q_g}$
= $\frac{T_e}{(T_c - T_e)} \cdot \frac{(T_g - T_a)}{T_g}$ (2.24)

2.3 COMPUTER PROGRAMME PACKAGE:

A computer programme has been developed to evaluate the C.O.P. of the single-stage-vapour-absorption-system, based on the thermodynamic analysis of the system described in section 2.2. Complete listing of this computer programme along with the description of the various input parameters and the execution commands on DECSYSTEM-1090 is given in Appendix-4.

The programme also includes the evaluation of pressure, temperature, concentration of LiBr-H₂O solution, enthalpy, mass flow rates of refrigerant and solution along with heat flow rates at various state points. The flow chart for the analysis is given in Fig. (2.2).

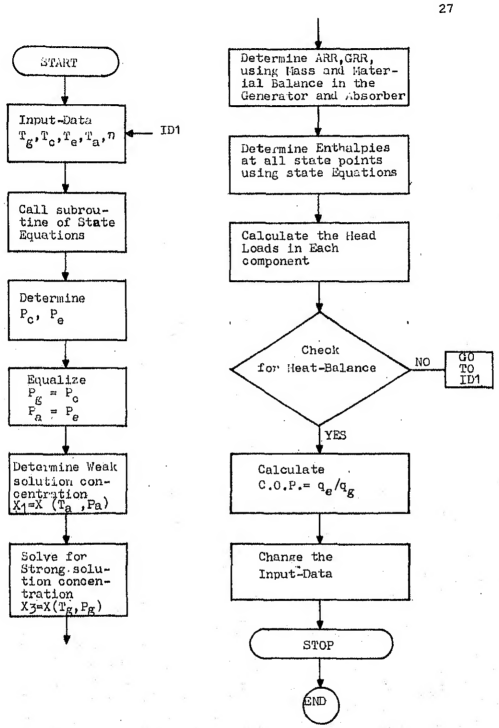


FIG. 2.2 FLOW CHART FOR A SINGLE-STAGE-VAPOUR-ABSORPTION-SYSTEM.

CHAPTER-III

ANALYSIS AND COMPUTER MODELLING OF DOUBLE-STAGE-VAPOUR-ABSORPTION-SYSTEM

3.1 DESCRIPTION OF THE SYSTEM :

Fig. (1.4) shows schematic diagram of the proposed double-stage-vapour-absorption-system. It operates with five different temperature level zones and three distinct pressure zones, as following:

The temperature levels are those of

- (i) generator-I, (ii) generator-II, (iii) condenser,
- (iv) evaporator, (v) absorber

The pressure zones are :

- (i) the high pressure in the generator-I, determined by the saturation pressure corresponding to the refrigerant temperature in the generator-II. The zone indicates a pressure range of 50-70 KPa.
- (ii) the medium pressure in the condenser and generator-II, in the range of 3-5 KPa. It is the saturation pressure corresponding to the condenser-temperature.
- (iii) the low pressure prevailing in the evaporator and absorber, is determined by the saturation pressure corresponding to the evaporator-temperature. It ranges from 0.7 to 1 KPa.

Heat is supplied from the heat source (solar, bio-gas, hot-water, hot-air, industrial waste-heat etc.) to the generator-I for the system operation. This heats the weak solution of LiBr-H₂O, pumped from the absorber into the generator-I, through heat exchangers-I and II. A percentage of the refrigerant i.e. water evaporates out of this weak solution and gets condensed in the generator-II. The remaining LiBr-H2O solution in generator-I passes through heat exchanger-II and a pressurereducing valve into the generator-II, where it receives latent heat from the refrigerant vapour evaporated out from generator-I. Some more percentage of refrigerant water from LiBr-H20 solution evaporates out and passes through the condenser. This evaporated water mixes with already condensed water coming out of the generator-II. Thus, the total amount of liquid refrigerant i.e. water leaving the condenser is the sum of the refrigerant originating from generators-I and II.

The refrigerant liquid from the condenser passes through an expansion valve, where its temperature and pressure lower down to the evaporator-temperature and pressure. In the evaporator, the refrigerant gets evaporated due to the latent heat of vaporisation, absorbed from the surroundings. This causes the cooling of space.

The cold vapours of the refrigerant, coming out of the evaporator are then dissolved in the strong solution of LiBr-H₂O,

collected in the absorber from the generator-II through the heat exchanger-I. A part of the heat of the strong solution of LiBr-H₂O coming out of generator-II is transferred to the weak solution of LiBr-H₂O, pumped from the absorber. Also, when the strong solution of LiBr-H₂O coming out of generator-II, mixes in the absorber with the low temperature and pressure water-vapour, coming out of the evaporator, it causes exothermic reaction and the heat so generated has to be rejected to the atmosphere by cooling the absorber. If the process of mixing is allowed to take place adiabatically (i.e. without rejecting the heat of absorption to the atmosphere), the temperature of the LiBr-H₂O solution will rise in the absorber and, eventually, the absorption of low temperature and pressure water-vapour would not take place.

Once the strong LiBr-H₂O solution mixes with the low temperature and pressure water-vapour in the absorber, the solution becomes weak and it is pumped out into the generator-I, as mentioned earlier. The pumping process elevates the pressure of the solution. Thus, the double-stage-vapour-absorption-cycle is completed.

3.2 THERMODYNAMIC ANALYSIS OF THE SYSTEM :

The thermodynamic analysis of the double-stage-vapourabsorption system is carried out to evaluate the system performance and other related parameters, such as pressure, temperature, concentration of solution, enthalpy, mass flow rate, heat flow rate at various state points. The assumptions for the analysis are the same as those for the single-stageabsorption-system given in Sec.(2.2). The details of this analysis are as following:

Condenser pressure ' P_c ', evaporator pressure ' P_e ', generator-II pressure ' P_g ', absorber pressure ' P_a ' and the concentration of LiBr- H_2 O solution, leaving the absorber ' X_6 ' are the parameters which can be determined by using the same procedures as were adopted for the single-stage-vapourabsorption-system, described in Section (2.2).

In generator-II, the pressure 'Pg2' is now known. Also, the source-temperature for heating the solution in generator-I and the solution temperature in this generator 'Tg1' are known. Hence, in both the components viz generator-I and generator-II, one property only is known for the solution in each component. The thermodynamic principles, however, require the prior knowledge of two independent properties, in order to define the state of solution uniquely. But the determination of second property for the solution in both generator-I and generator-II, presents a problem, which is solved by adopting the following procedure:

First of all, the concentration of LiBr-H $_2$ 0 solution 'X $_{12}$ ' leaving the generator-II is assumed. Since, the 'P $_{g2}$ '

in the generator-II is already known, P_{g2} and X_{g2} i.e. X_{12} are the two properties which fix the thermodynamic state of the solution. Therefore, the solution temperature leaving generator-II can be determined by using Appendix-2. Assuming a terminal temperature difference of P_{g2} and the refrigerant vapour coming from generator-I P_{g2} and the solution temperature P_{g2} , for heat-transfer between the two fluids, P_{g2} can also be determined. Using Appendix-2, the vapour pressure of the refrigerant corresponding to temperature P_{g2} can be directly read. This pressure P_{g2} corresponds to the pressure in generator-I. Since P_{g2} is already known, the concentration of the LiBr-H₂O solution P_{g2} already known, generator-I can be evaluated with the help of Appendix-2.

We now use the mass, material (LiBr), heat and energy balances in the following components of the system. The details of which are presented below:

(a) Generator-I

Mass balance :

$$\dot{m}_8 = \dot{m}_1 + \dot{m}_{10} \tag{3.1}$$

Material balance (LiBr Conservation) :

$$\dot{m}_8 X_8 = \dot{m}_{10} X_{10}$$
 (3.2)

(b) Generator-II

Mass balance :

$$\dot{m}_{11} = \dot{m}_{12} + \dot{m}_{9}$$
 (3.3)

Material balance :

$$\dot{m}_{11} X_{11} = \dot{m}_{12} X_{12}$$
 (3.4)

Energy balance :

$$\dot{m}_1(H_1-H_2) = \dot{m}_{12} H_{12} + \dot{m}_9 H_9 - \dot{m}_{11} H_{11}$$
 (3.5)

(c) <u>Heat-Exchanger-I</u>

Heat balance :

$$\dot{m}_6(H_7-H_6) = \dot{m}_{12}(H_{12}-H_{13})$$
 (3.6)

Heat exchanger effectiveness :

$$n_1 = \frac{\text{Useful enthalpy drop}}{\text{Maximum possible enthalpy drop}}$$

$$= \frac{H_{12} - H_{13}}{H_{12} - H_{6}}$$
(3.7)

(d) Heat-Exchanger-II

Heat balance :

$$m_7(H_8-H_7) = m_{10}(H_{10}-H_{11})$$
 (3.8)

Heat exchanger effectiveness

$$n_2 = \frac{H_{10} - H_{11}}{H_{10} - H_7} \tag{3.9}$$

(e) Absorber

Mass balance :

$$\dot{m}_6 = \dot{m}_5 + \dot{m}_{13} \tag{3.10}$$

Material balance :

$$\dot{m}_6 X_6 = \dot{m}_{13} X_{13}$$
 (3.11)

(f) Condenser

Mass balance :

$$\dot{m}_9 + \dot{m}_2 = \dot{m}_3$$
 (3.12)

In expressions (3.1) to (3.12), the parameters

 1 H₁, 1 H₂, 1 H₅, 1 H₆, 1 H₉, 1 H₁₀, 1 H₁₂, 1 X₆, 1 M₁ and 1 M₂ are known. When these known parameters are substituted in the above expressions, they generate a set of six independent equations in six unknown variables, which are as following:

$$\dot{m}_1 (H_1 - H_2) = \dot{m}_{12} H_{12} + \dot{m}_9 H_9 - m_{11} H_{11}$$
 (3.13)

$$\dot{m}_7 (H_8 - H_7) = \dot{m}_{10} (H_{10} - H_{11})$$
 (3.14)

$$\dot{m}_{12} (H_{12} - H_{13}) = \dot{m}_7 (H_7 - H_6)$$
 (3.15)

$$\dot{m}_{11} = \dot{m}_9 + \dot{m}_{12}$$
 (3.16)

$$\dot{m}_3 = \dot{m}_1 + \dot{m}_9 \tag{3.17}$$

$$H_{10}-H_{11} = n_2 (H_{10}-H_7)$$
 (3.18)

The unknown variables in the expressions (3.13) to (3.18) are : \dot{m}_1 , \dot{m}_9 , \dot{m}_{12} , \dot{H}_7 , \dot{H}_8 and \dot{H}_{11}

Simultaneous solution of equations (3.13) to (3.18) is then obtained with the help of MATRIX INVERSION METHOD. This solution yields all the unknown parameters.

Using expressions (3.1) and (3.2), the Generator-Recirculation-Ratio, 'GRR 'defined in Section (2.2), for the present system is given by,

GRR = Mass of LiBr-H₂O solution, leaving the generator-I
Mass of water-vapour driven off from the generator-I

$$= \frac{\mathring{m}_{10}}{\mathring{m}_{1}}$$

$$X = \frac{x_8}{x_{10} - x_8}$$

$$\therefore \frac{\dot{m}_{10}}{\dot{m}_{1}} = \frac{x_{8}}{x_{10} - x_{8}} \tag{3.19}$$

All the variables in the above expression are known except \dot{m}_{10} , which can be easily determined.

Expression (3.4) for LiBr balance in generator-II is represented as :

$$\dot{m}_{11} X_{11} = \dot{m}_{12} X_{12}$$
 (3.20)

Since,

$$\dot{m}_{11} = \dot{m}_{10}$$
,

and,

 $X_{11} = X_{10}$, the expression (5.20) reduces to ,

$$\dot{m}_{10} = \dot{m}_{12} \times 12$$

or,

The concentration X_{12} of the LiBr- H_2 O solution is given by,

$$X_{12} = \frac{\dot{m}_{10} X_{10}}{\dot{m}_{12}} \tag{3.21}$$

All the parameters on the R.H.S. of the above expression are known and, hence, using (3.21), X_{12} can be determined.

This calculated value of X_{12} is then compared with the value of this parameter, assumed to be known initially. If they are not in close agreement with each other, then a different value of X_{12} is assumed and the above procedure is repeated till the difference between the assumed and calculate value of X_{12} becomes negligible.

The system performance, represented by C.O.P. is then obtained as :

C.O.P. = Refrigerating effect in the evaporator
Heat supplied in the generator-I

$$= \frac{\dot{m}_{4} (H_{5} - H_{4})}{\dot{m}_{10} H_{10} + \dot{m}_{1} H_{1} - \dot{m}_{8} H_{8}}$$
 (3.22)

The overall heat-balance of the complete system is performed, which corroborates the correctness of the solution-method presented above. The heat balance equations for various components are:

(i) Heat addition in generator-I:

$$q_{g1} = \dot{m}_{10} H_{10} + \dot{m}_{1} H_{1} - \dot{m}_{8} H_{8}$$
 (3.23)

(ii) Heat transfer in generator-II:

$$q_{g2} = \dot{m}_1 (H_1 - H_2)$$

$$= \dot{m}_{12} H_{12} + \dot{m}_9 H_9 - \dot{m}_{11} H_{11}$$
 (3.24)

(111) Heat rejection in condenser :

$$q_c = \mathring{m}_9 + \mathring{m}_2 + \mathring{m}_2 + \mathring{m}_3 + \mathring{m}_3$$
 (3.25)

(iv) Heat extracted in the evaporator:

$$q_e = \dot{m}_4 (H_5 - H_4)$$
 (3.26)

(v) Heat rejection in the absorber :

$$q_a = \dot{m}_{13} + \dot{m}_{5} + \dot{m}_{5} + \dot{m}_{6} + \dot{m}_{6}$$
 (3.27)

The final heat balance is given by, Total heat inflow into the system

= Total heat outflow from the system or,

$$q_e + q_{g1} = q_c + q_a$$
 (3.28)

This completes the thermodynamic analysis of the proposed double-stage-vapour-absorption-system.

3.3 COMPUTER MODELLING :

A computer programme package given in Appendix-5 has been developed on the basis of the thermodynamic analysis given above. The complete logic, which forms the basis for the development of the programme has been mentioned in the flow-chart, given in Fig. (3.1). Various subroutines developed to solve state equations of LiBr-H₂O solution and refrigerant along with other parameters are given in Section (3.3.1).

Newton-Raphson's Numerical Technique (Sec. 3.3.2) has been used to solve the non-linear algebric equations in the problem. Subroutines, given by Numerical Aløgorithm Group (NAG), are employed to solve the simultaneous linear equations. The programme has been made very general to encompass a class of similar problems, which can be solved by simply changing input variables in the parent programme.

3.3.1 Subroutines for solving state equations :

The various subroutines developed for solving the thermodynamic state equations (Appendix-1) for the LiBr-H₂O solution and refrigerant are given below:

(i) subroutine for the pressure of saturated refrigerant vapour when its temperature is known:

$$P = P (TR) (3.29)$$

(ii) subroutine for the concentration of the solution when its temperature and refrigerant temperatures are known:

$$X = X (TS, TR)$$
 (3.30)

(iii) subroutines for the concentration of the solution when refrigerant temperature and pressure are known:

$$X = X (P, TR)$$
 (3.31)

(iv) subroutine for enthalpy of solution, when solution temperature and concentration are known.

$$H = H (TS, X)$$
 (3.32)

(v) subroutine for the solution temperature when its enthalpy and concentration are known.

$$TS = TS (H, X) (3.33)$$

3.3.2 Algorithem, Newton-Raphson Technique:

The Newton-Raphson iterative technique is used for solving higher-order, non-linear algebric equations, as follows:

A function f (X), which is continuous and has a continuous derivative is given. Also given is a starting value $\rm X_{O}$ (the concentration of LiBr-H₂O solution in the present case).

For $n = 1, 2, \dots$ until termination, do: compute $f'(X_n)$

If $f'(X_n) = 0$, signal and stop.

Else, compute

$$x_{n+1} = x_n - \frac{f(x_n)}{f'(x_n)}$$
 (3.34)

till difference between the values \mathbf{X}_{n+1} and \mathbf{X}_n tends to zero.

The termination criterion used for iteration:

After N steps (N assumed, fixed)

or,

$$|(X_{n+1} - X_n)| \leqslant \epsilon \quad (\epsilon > 0, \text{ given})$$

or,

$$|f(X_n)| \leqslant \alpha \quad (\alpha > 0 \text{ given })$$

Stop.

The flow-chart for the analysis is given in Fig. (3.1).

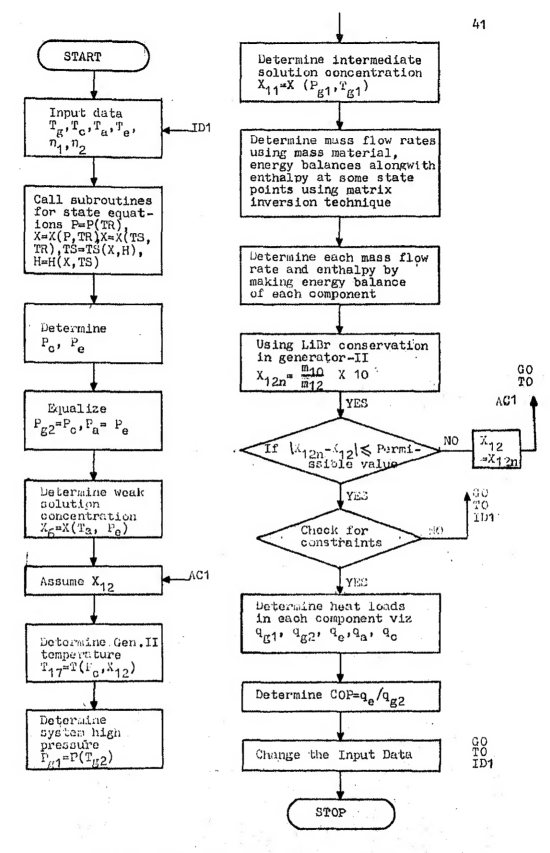


FIG.3.1 FLOW-CHART FOR COMPUTER MODELLING OF A DOUBLE-STAGE-VAPOUR-ABSORPTION-SYSTEM

CHAPTER- IV

ECONOMICS OF VAPOUR ABSORPTION AND VAPOUR-COMPRESSION SYSTEMS

OPERATIONAL COST :

A comparative cost-analysis is carried out to study the relative economics of the present system with its most competing alternative i.e. the-vapour-compression-system. As a case-study, the air-conditioning unit, installed at I.I.T. Kanpur which operates on a vapour-compression -cycle having R-22 as refrigerating fluid has been considered. The detailed data for the air-conditioning unit are presented in Appendix 6. The total cost of any of the above two system involves the following two costs:

- (i) first cost,
- (ii) running or operational cost.

It has been reported in Reference \(\frac{7}{20} \) 7 that the first cost of the two system is almost the same. We shall, therefore, compare only the running cost of the two systems.

The running cost of the vapour-compression-system consists of the following :

(a) Cost of the electrical energy used for pumping cooling-water in the condenser.

Since, the condenser heat load in both the systems vary, the electrical energy used will also be different for the two systems. This has been discussed in detail later.

(b) Cost of the electrical energy, required for circulating the chilled water in the evaporator.

As the cooling capacity of the evaporator in both the system is taken as the same, the electrical energy used for the purpose, will be the same for both the systems. Hence it contribution has not been considered.

(c) Cost of the electrical-energy consumed in the compressor.

Contribution of this cost is maximum in running the compression-system. Its inclusion for evaluating the total running cost of the system is, therefore a must.

Details of the method for obtaining the total running cost of the vapour-compression-system is presented as follows:

Capacities of the condenser pumps and compressors, used in the system and considered for the case study are reported in (Appendix-6).

Suppose,

 PWR_c = Total power of the compressors /KW_7

Total power requirement of the complete system will be given by,

Total power =
$$PWR_T = PWR_P + PWR_c$$
 (4.1)

Annual electrical energy consumption, considering continuous running of the system is then represented as,

$$(AEC)_{VCS} = PWR_T \times 24 \times 365$$
 _KW-hr_7 (4.2)

In the above expression, 'AEC' represents the 'ANNUAL ENERGY CONSUMPTION' and the subscript 'VCS' stands for the vapour-compression-system.

Let the cost of electrical energy be denoted by 'Cen' in _Ts/KW.hr_7.

The total cost involved in meeting the running cost of the system annually is, then, given by ,

$$(TRC)_{VCS} = (AEC)_{VCS} \times Cen$$
 (4.3)

where,

TRC represents the total running cost involved.

The total running-cost of the double-stage-vapour-absorption-system (DSVAS) is then obtained as following:

Factors, affecting the running cost of the system are :

- (1) Cost of the electrical energy, consumed for pumping cooling-water in the condenser.
- (2) Cost of the electrical energy, required for the solution-pump to raise the pressure of the LiBr-H₂O solution from the absorber pressure to generator pressure.

The cost-element, given in (1) is determined as follows:

The mass flow rate of the cooling-water is determined by using the relation :

$$q_{C} = \dot{m}_{W} \times C_{PW} \Delta T \qquad (4.4)$$

In the above expression, 'q_c', 'C_{PW} and ' Δ T'are known parameters.'q_c' is condenser heat load obtained numerically using the computer programme given in Appendix -5; C_{PW} the specific heat of water, is taken from the table given in $\int 18_7$; Δ T is the allowable temperature rise in the condenser which has been assumed as 6° C in our case (Appendix-6). Hence the unknown parameter \dot{m}_{W} is determined from the expression (4.4).

The volume flow rate of cooling-water i.e. C through the condenser is then given by,

$$Q = \frac{\dot{m}_W}{\rho_W} \tag{4.5}$$

The pressure-drop $^{\bullet} \triangle P$ $^{\bullet}$ through the condenser tube is calculated as following :

$$p = \frac{fL V_W^2}{2 d_i} \rho_W \tag{4.6}$$

In the above expression, L', V'_w and f' represent the length of tubes, velocity of water and the friction factor respectively. Details of the methods for their determination are discussed below:

Length of condenser tubes :

The condenser heat load q_c is given by the relation, $q_c = U_o \times A_o \times LHTD$

$$\therefore A_o = \frac{q_c}{U_o \times LMTD}$$
 (4.7)

$$LMTD = \frac{\Delta T_{i} - \Delta T_{o}}{\ln \frac{\Delta T_{i}}{\Delta T_{o}}}$$
 (4.8)

The overall heat transfer coefficient U is given by

$$U_{o} = \frac{1}{R_{o} + R_{t} + R_{ff} + R_{i}}$$

$$= \frac{1}{\frac{1}{h_{con}} + \frac{b d_{o}}{R_{m} d_{m}} + \frac{d_{o}}{h_{ff} d_{i}} + \frac{d_{o}}{h_{w}} \frac{1}{d_{i}}}$$
(4.9)

Where,

= 0.725
$$\left(\frac{g \rho_W^2 H_{fg} K_W^3}{\mu \text{ t N d}_0}\right)^{0.25}$$
 (4.10)

K, = conductivity of water

 μ = viscosity of water

N = number of tubes in one vertical row, assumed to be 3 as explained in $\boxed{79.7}$

 d_{o} = outer dia of tube, assumed to be 18 mm (Appendix-6)

 $H_{fg} = Latent heat of vaporisation of water$

t = temperature difference between the vapour and surface of the tube (initially assumed for iteration purpose as 5°C)

 $h_{ff} =$ fouling factor taken

= \cdot 000176 (m^2 K/W), \triangle Reference-19_7

 d_1 = inner dia of tube, assumed to be 16 mm (Appendix-6)

d_m = mean diameter

$$= \frac{d_0 + d_1}{2}$$

b = thickness of tube

$$= \frac{d_0 - d_1}{2}$$

 K_{m} = conductivity of tube material

 $h_{\rm W}$ = water-side heat transfer coefficient * After substituting all the parameters in expression (4.9), the overall heat transfer coefficient $U_{\rm O}$ is obtained. How all the terms on the R.H.S. of expression (4.7) are known. Hence $A_{\rm O}$ can be determined from the above expression. Also, the condenser load is given by ,

$$q_c = h_{con} \times A_o \times \Delta t \tag{4.11}$$

$$\therefore \Delta t = \frac{q_c}{h_{con} \times A_o}$$
 (4.12)

If the calculated value of t does not match with its assumed value in expression (4.10), new value of Δ t is assumed and the entire procedure to calculate Λ_0 is repeated. The calculations are terminated, when the difference between the assumed and the calculated value of t is negligible.

where, n and m are exponents. The constant C and exponents in the above expression are given by,

$$\frac{h_W d_i}{K_W} = 0.023 \left(\frac{V d_i \rho_W}{\mu} \right)^{0.8} x \left(\frac{C_p}{K} \right)^{0.8} / \text{Reference-9_7}$$

^{*} The expression for the heat transfer coefficient for fluids flowing inside tubes, i.e. water in this case, is of the form

The total outer area of tubes, is also given by,

$$A_{O} = \pi \times d_{O} \times L \tag{4.13}$$

where,

L is the total length of tubes.

$$L = \frac{A_0}{\pi d_0} \qquad (4.14)$$

Thus, the required total length of the tubes, used in the condenser, is known.

Velocity of water in tube :

The velocity of water $\,{\rm V}_{\rm W}\,\,$ is obtained from the relation

$$V_{W} = \frac{Q}{\frac{\pi}{4} \times d_{i}^{2}} \tag{4.15}$$

where, Q is the volume flow rate of water, obtained in expression (4.5).

Friction-factor:

The friction-factor is given by $\sqrt{17}$,

$$f = \frac{1}{1.14 + 2 \log \frac{d_i}{\epsilon} - 2 \log / 1 + \frac{9.3}{Re(\epsilon/d_i) f} / 7^2}$$

(4.16)

where.

' ϵ ' is absolute roughness of pipe material given in Appendix -7.

Re is given by

$$Re = \frac{V_W d_i \rho_W}{\mu}$$
 (4.17)

Since, the equation (4.16) is implicit in 'f', Newton-Raphson numerical technique (Sec. 3.4) is adopted to solve this equation and to obtain the value of 'f'.

The pressure drop $^{1}\Delta p^{1}$ can then be easily obtained, using expression (4.6).

The power required for pumping is then given by,

$$(PWR)_{cond} = \Delta p \times Q \qquad (4.18)$$

In the above expression, the subscript cond stands for condenser .

The annual cost of the electrical energy represented as $(TRC)_{cond}$ is then expressed as :

$$(TRC)_{cond} = (PWR)_{cond} \times 24 \times 365 \times C_{en}$$
 (4.20)
where,

'C is the cost of electrical energy in Rs./KW-hr.

The second running cost-element i.e. the cost of electrical energy required for the solution pump to raise the pressure. Referring to Fig. (1.4), the pressure against which

the the solution pump has to work, comprises of the following pressure drops :

- (i) pressure difference between the generator and absorber (Δp_1),
- (ii) pressure drop in solution heat-exchanger-I (Δp_2) and
- (iii) pressure drop in solution heat-exchanger-II (△p3)
- (iv) pressure drop in pipe length (Δp_4) P_4 is determined by using the relation

$$P_1 = P_{g1} - P_a$$
 (4.21)

where,

Pg1 = generator-I pressure,

P_a = absorber pressure

 P_{g1} and P_{a} are known from Appendix -5

 Δ p₂ and Δ p₃ are evaluated by following the same procedure as adopted for obtaining Δ p of expression (4.1) for condenser cooling-water. The properties of LiBr-H₂O solution, given in Reference [26.7], are however used now, instead of the properties of water.

The pressure drop \triangle p₄ in pipe length can be neglected as it is negligible as compared to above pressuré drops.



The total pressure drop which is to be supplied by the solution pump is therefore, given as:

$$\Delta P_{T} = \Delta P_{1} + \Delta P_{2} + \Delta P_{3} \tag{4.22}$$

From the knowledge of P_{T} , the power required for solution pump (PWR)_{\mathrm{SP}} can be determined as :

$$PWR_{SP} = \Delta P_{T} \times Q_{S}$$
 (4.23)

where,

 $Q_{\rm S}$ = solution flow rate, obtained from Appendix-5.

Total running cost for solution pump can be expressed as :

$$(TRC)_{SP} = (PWR)_{SP} \times 365 \times 24 \times Cen (4.24)$$

The total running cost, incurred annually for the complete system operation of double-stage-vapour-absorption-system is given by,

$$(TRC)_{DSVAS} = (TRC)_{cond} + (TRC)_{SP}$$
 (4.25)

here,

 ${\rm (TRC)}_{\rm DSVAS}$ stands for the total running cost of the double-stage-vapour-absorption-system, incurred annually.

A complete computer-programme on DEC-1090, given in Appendix-8, has been developed to solve the above problem and obtain the total running-cost of the two systems. From the programme, we obtain

(a)
$$(TRC)_{DSVAS} = Rs. 3,73,350.00$$

(b)
$$(TRC)_{VCS} = Rs. 8,08,000.00$$

Therefore the annual saving in running cost with the double-stage vapour-absorption-system as compared to vapour-compression-system is given by,

(c) Annual saving =
$$(TRC)_{VCS}$$
- $(TRC)_{DSVAS}$
= Rs. 4,34,650.00

or

- (d) Daily saving = Rs. 1,190.00
- (e) Percentage saving = 53.8 %.

CHAPTER-V

RESULTS AND DISCUSSION

absorption-system, using thermodynamic equations of state has been developed to determine the feasibility and economy of its operation. DEC-1090 computer with FORTRAN as coding language has been used for executing the programme. A numerical model of one-stage-vapour-absorption-system has also been developed to compare its relative performance with that of a double-stage-system.

The results of the above analyses have been presented in Tables 1.11. On the basis of the data obtained from these tables, figures (5.1 - 5.11) have been drawn.

Figures (5.2-5.4) show the variation of C.O.P. with generator-I temperature for both the systems. Also, the minimum temperatures at which double-stage-absorption-system starts operating for different cooling water temperatures, have been shown. It is observed that this system starts working at a minimum temperature of 70°C, when cooling water temperature is 20°C and at about 90°C for cooling water-temperature of 35°C.

An interesting result is observed from Fig. (5.3). The crystallization of LiBr-H₂O solution begins at around 91°C for a single-stage-system. But, this is the range of temperature, at which a double-stage-vapour-absorption-system starts working. This establishes the fact that when the single-stage-vapour-absorption-system stops functioning, the double-stage-system starts operating. Figs.(5.2-5.4) show that the maximum value of C.O.P. obtained in the double-stage-system is almost twice the value of the corresponding single-stage-system (~1.7 as compared to ~0.85). This is due to the fact that about 80-90 % of the energy input in the generator-I is recovered as the heat of condensation in generator-II, to further drive off the refrigerant vapour, which otherwise would have been wasted in the condenser.

As C.O.P. is the index of effectiveness of the system, it depicts that a double-stage-vapour-absorption-system has almost half the energy consumption per unit cooling achieved; in comparison to a single-stage-system.

The variation of generator-I temperature V/S C.O.P. in a double-stage-system shown in Fig. (5.1) reveals that, upto a certain increase in generator-I temperature, C.O.P. also increases and afterwards it starts decreasing. The optimum value of generator-I - temperature for maximum C.O.P. can, therefore, be obtained. The optimum value of generator-I

temperature is different for different condensing and evaporating temperatures. From Figure (5.1), it is obvious that for 20°C as condenser-temperature and 5°C as evaporator temperature, the optimum value of generator-I-temperature is 79°C and for 25°C as condenser temperature and 5°C as evaporator temperature, the optimum generator-I-temperature for maximum C.O.P. is 88°C. Maximum C.O.P. is 1.785 in this case.

C.O.P. of the double-stage-vapour-absorption-system falls with the increase in the absorber-temperature. Reduction in C.O.P. is less, initially, but after a certain value of the absorber-temperature, the curve becomes quite steep. This is due to the fact that at higher absorber-temperature, absorption capacity of LiBr-H₂O solution to absorb the refrigerant vapour coming from the evaporator decreases. The reduction in the absorption capacity requires more solution to flow for absorbing a given amount of refrigerant. Therefore, more heat is needed in generator-I to extract the refrigerant from the solution.

Figures (5.5-5.6) show that with the increase in condenser-temperature, C.O.P. decreases. However, the rate of reduction is less than that of reduction in C.O.P. with absorber-temperature Fig. (5.8). It shows that variation in C.O.P. is much more sensitive to the absorber-temperature than the condenser-temperature.

At high condensing temperatures such as at atmospheric temperature of water or air, the double-stage-system performs effeciently (Fig. 5.6). Hence, the cooling of condenser can be performed at atmospheric temperature of water or air. The need of a cooling tower, which is generally used to reduce the cooling-water temperature, can therefore be eliminated.

The effectiveness of the heat exchangers-I and II have a significant effect on the C.O.P. With the increase in the effectiveness, the C.O.P. increases (Fig. 5.10) and varies between 1.325 to 1.778 for the corresponding effectiveness 0.5 to 1.0 of heat exchanger-I and II.

Figure 5.9 shows that generator-I heat load first decreases rapidly with the increase in generator-I-temperature, but afterwards it increases slightly. The inversion point of this curve corresponds to the maximum C.O.P. of the double-stage-system. This amounts to conclude that at the optimum generator-I-temperature, the heat load required to be supplied in generator-I is minimum.

Since the double-stage-system uses low grade heat energy for its operation as compared to high grade mechanical energy used by vapour-compression-system, the former has the potential advantages over the later, as far as the operational costsof the

systems are concerned. For the same system capacity of 550 tons that has been studied in the present work, as an case study, the yearly saving achieved in the running cost of a double-stage-vapour-absorption-system is approximately 4.35 lakhs over the vapour-compression system.

Numbers at top of each column in the tables (1-4) denote the following parameters:

```
1. Serial No;
```

- 2. Generator-I temperature (°C);
- Generator-II temperature (OC);
- 4. Generator-I load (KW);
- 5. Evaporator load (KW);
- 6. Generator-II load (KW);
- 7. Weak-solution concentration (Kg LiBr/Kg of solution);
- 8. Intermediate-solution concentration (Kg LiBr/Kg of solution);
- 9. Strong solution concentration (Kg LiBr/Kg of solution);
- 10. C.O.P., Double-stage;
- 11. C.O.P., Single-stage;
- 12: C.O.P., Ideal, absorption cycle
- Cry. Crystallization
- ONP Operation not possible

Number at top of each column in the Tables (5-11) denote the following parameters:

- 1 Serial No.
- Variable indicated in the table column
- 3 Generator-II temperature (OC);

```
4 C.O.P. (Double-stage)
```

- 5 C.O.P. (Single-stage)
- 6 C.O.P. (Ideal-cycle)

For the Table. 12 the numbers at the top of column denote:

- 1 Serial No.
- 2 Heat-Exchanger Effectiveness
- 3 C.O.P. (Double-stage)
- 4 C.O.P. (Single-stage)

TABLE -1

Effect of Generator-I Temperature on operating parameters

		E-1)	$(T_c = 20^{\circ}C,$		$T_a = 20^{0}$ C	့	Te= 5°C	ر ن	$n_1 = n_2 =$	$n_1 = n_2 = 0.90$	
1	2	2	4	5	9	7	ω	o,	10	11	12
1	52.0	31								.8395	
	58.0									.8349	
	64.0				#5 					.8514	
	70.07	47.1	9.985	17.747	9.064	45.76	48.19	50,60	1.77	.8295	2,702
	75.0	49.2	9.952	17.747	9.081	45.76	48.75	51.70	1.780	.8276	2.839
	0.97	51.0	3,945	17.747	6.097	45.76	49.20	52.80	1.785	Cry.	2.974
	0.67	52.8	6,945	17.747	9.111	45.76	97.64	79.64	53.90	1.786	3.107
	82.0	55.0	9,6,6	17.747	9.122	45.76	50.14	54.90	1.783		5.237
	85,0	57.0	9.954	17.747	9.132	45.76	50,60	55.90	1.781		3.365
	0.58	59.0	9,965	17.747	9.143	45.76	51.02	56.90	1.779		5.491
	91.0	9.09	6.975	17.74.7	9.149	45.76	51.55	57,80	1.778		3.615
	0.476	62.6	766.5	17.747	9.161	45.76	51.89	50.80	1.776		5.737
	97.0	64.7	10,008	17.747	9.167	65.76	52.35	59.70	1.772		5.857
	100.0	4.99	10,025	77.747	9.176	92.54	52.77	50.50	1.770		3.975

TABLE-2

Effect of Generator-I Temperature on operating parameters

		12	1.396	1.483	1.596	1.706	1.815	1.922	2,026	2,427	2,524	2,618	2.711	2.803	2,893	2.981
		17	.8277	.8271	8247	.8216	.8210	.8196	.8180	Cry						
parameters	2 = 0.90)	10				1.695	1.724	1.740	1.747	1.750	1.752	1.752	1.751	1.749	1.746	1.742
operating p	$n_1=n_2$	6				48.20	04, 64	50,50	51.60	52.70	53.70	54.70	55.70	56.70	57.70	58.60
e e e	, D	8				47.06	47.60	48.17	48.69	49.16	89.67	50.17	50.62	51.03	51.40	51.86
TABLE-2 of Generator-I Temperature	$T_e = 5^{\circ}C$	7				45.76	45,76	45,76	45.76	45.76	45.76	45.76	45.76	92°24	45.76	45.76
TAI	, 20°05	9	÷		- ONP	6.067	9.087	660.6	9.109	9.120	9.128	9.135	9.14	9.153	9,163	9.168
Effect of G	$T_a = 20$	5				17.594	17.594	17,594	17.594	17,594	17,594	17.594	17.594	17,594	17,594	17,594
<u>급</u>	25°C, 2	4				10.378	10.204	10,109	10.069	10.054	10.041	10.040	10.047	10.059	10.074	10.086
	$T_e = 25$	3				8.84	50.7	52.7	54.5	9.95	58.5	4.09	62.1	64.1	66.1	0.89
	, As	2	52.0	58.0	-0.49	70.0	75.0	76.0	79.0	82.0	85.0	88	91.0	0.46	97.0	100.0
		~	\ \ <u>\</u>	Ŋ	M	7	ľ	٥	7	80	6	10	<u></u>	12	13	14

TABLE -3

Effect of Generator-I Temperature on operating parameters

3.7560 1.106 3.7560 1.106 3.882 49.34 49.56 49.30 0.666 .8080 1.786 3.026 49.34 49.85 50.40 1.438 .8052 1.864 3.026 49.34 50.43 51.40 1.600 Cry. 1.941 3.079 49.34 51.45 55.40 1.676 2.091 3.079 49.34 51.90 54.40 1.676 2.091 3.097 49.34 52.80 55.30 1.698 2.236 3.131 49.34 52.80 55.30 1.704 2.307
49.34 49.56 49.30 0.666 49.34 49.85 50.40 1.438 49.34 50.43 51.40 1.600 49.34 50.96 52.40 1.652 49.34 51.45 53.40 1.676 49.34 51.45 53.40 1.696 49.34 51.90 54.40 1.696 49.34 52.42 55.30 1.696 49.34 52.80 56.30 1.701 49.34 52.25 57.20 1.704
.8086 .8115 .8115 .9032 .90,34 49.85 60.40 1.438 .8052 .49,34 50.43 51.40 1.600 Cry49,34 50.96 52.40 1.652 .49,34 51.90 54.40 1.676 .49,34 52.80 55.30 1.698 .49,34 52.80 56.30 1.701 .49,34 52.80 56.30 1.704
.8115 49.54 49.56 49.50 0.666 .8080 49.34 49.85 50.40 1.438 .8052 49.34 50.45 51.40 1.600 Cry. 49.34 51.45 55.40 1.676 49.34 51.90 54.40 1.696 49.34 52.82 55.30 1.698 49.34 52.82 55.30 1.701
49.34 49.56 49.30 0.666 .8080 49.34 49.85 50.40 1.438 .8052 49.34 50.43 51.40 1.600 Cry. 49.34 50.96 52.40 1.652 49.34 51.45 53.40 1.676 49.34 51.45 55.40 1.696 49.34 51.90 54.40 1.696 49.34 52.42 55.30 1.698 49.34 52.80 56.30 1.701 49.34 52.25 57.20 1.704
49.34 49.56 49.30 0.666 .8080 49.34 49.85 50.40 1.438 .8052 49.34 50.43 51.40 1.600 Crry. 49.34 50.96 52.40 1.652 49.34 51.45 53.40 1.676 49.34 51.90 54.40 1.696 49.34 51.90 54.40 1.696 49.34 52.42 55.30 1.696 49.34 52.42 55.30 1.696 49.34 52.25 57.20 1.704
49.34 49.85 50.40 1.438 .8052 49.34 50.43 51.40 1.600 cry. 49.34 51.45 52.40 1.676 49.34 51.45 53.40 1.676 49.34 51.90 54.40 1.690 49.34 52.42 55.30 1.696 49.34 52.42 55.30 1.701 49.34 52.80 56.30 1.701 49.34 52.25 57.20 1.704
49.3450.4351.401.600Cry.49.3450.9652.401.65249.3451.4553.401.67649.3451.9054.401.69049.3452.4255.301.69849.3452.8056.301.70149.3453.2557.201.704
49.3450.9652.401.65249.3451.4553.401.67649.3451.9054.401.69049.3452.4255.301.69849.3452.8056.301.70149.3453.2557.201.704
49.3451.4553.401.67649.3451.9054.401.69049.3452.4255.301.69849.3452.8056.301.70149.3453.2557.201.704
49.34 51.90 54.40 1.690 49.34 52.42 55.30 1.698 49.34 52.80 56.30 1.701 49.34 53.25 57.20 1.704
49.34 52.42 55.30 1.698 49.34 52.80 56.30 1.701 49.34 53.25 57.20 1.704
49.34 52.80 56.30 1.701 49.34 53.25 57.20 1.704
49.34 53.25 57.20 1.704

TABLE-4

Effect of Generator-I Temperature on operating parameters

()	12					1.525	1.599	1.672	1.744	7.015	1.955	2.021	2.087	2.152
$n_1 = n_2 = 0.90$	11	.2897	.7844	.7968	. 7986	. 7976	.7967	Cry.						
n_{7}	10						97.0	1.231	1.514	1.593	1.535	1.652	1,000	1.675
$T_e = S^0C$,	6						50.00	51.00	52.00	55.00	53.90	54.90	55,30	56.70
6-1 11	8						50.20	50.70	51.22	51.70	52.25	52.65	53.13	55.57
	7						50.39	50.39	50.39	50.39	50.39	50.39	50.39	50.39
, 30°C ;	9	3-				ONP	8.678	8,762	8.964	9.018	140.6	790.6	6.077	680.6
 	5						17.328	17.328	17.328	17.328	17,328	17.328	17,328	17.528
35°C,	4	. 0					17.526	14.076	11.146	10.880	10.589	10.489	10,405	10,359
O	5	,					62.0	64.1	9.00	66.5	70.07	7.0	73.7	75.6
	2	0.49	70.0	76.0	82.0	88.0	91.0	0.46	0.79	100.0	103.0	106.0	109.0	112.0
	-	-	2	M	7	īV	Q	7	Ø	σ	10	7	12	12

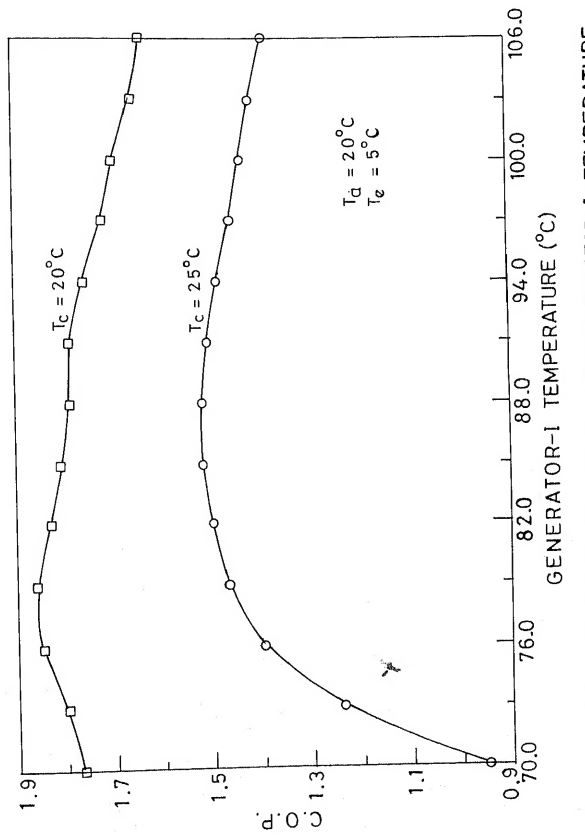


FIG. 5.1 VARIATION OF C.O.P. WITH GENERATOR-I TEMPERATURE.

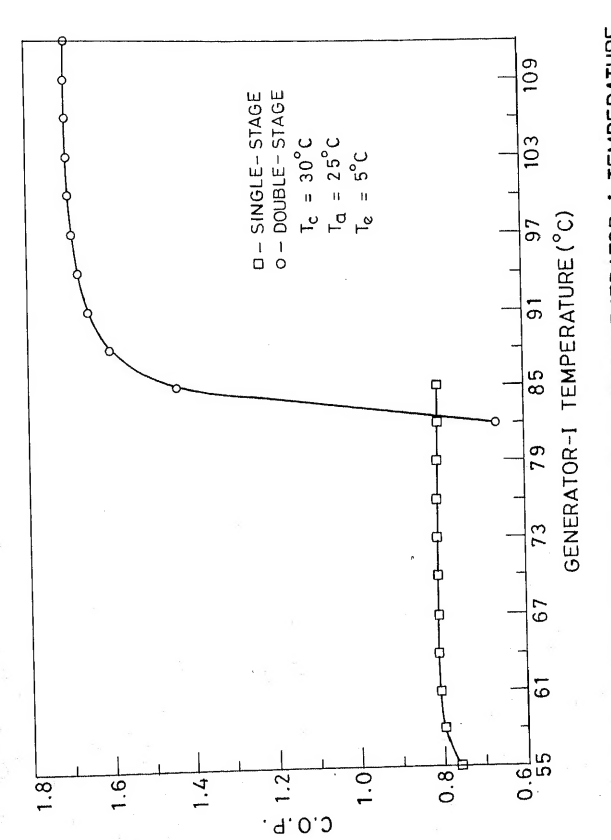
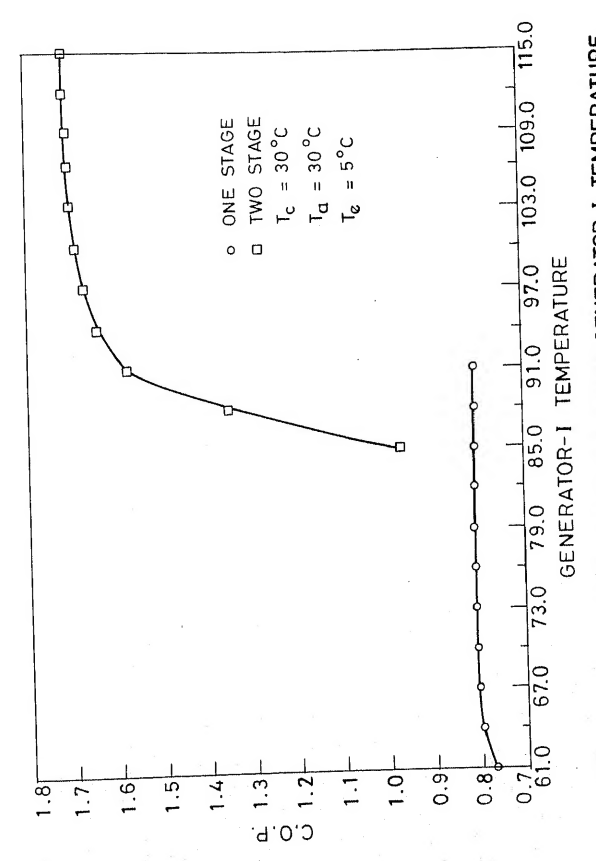


FIG.5.2 VARIATION OF C.O. P. WITH GENERATOR-I TEMPERATURE.



VARIATION OF C.O.P. WITH GENERATOR-I TEMPERATURE. FIG. 5.3

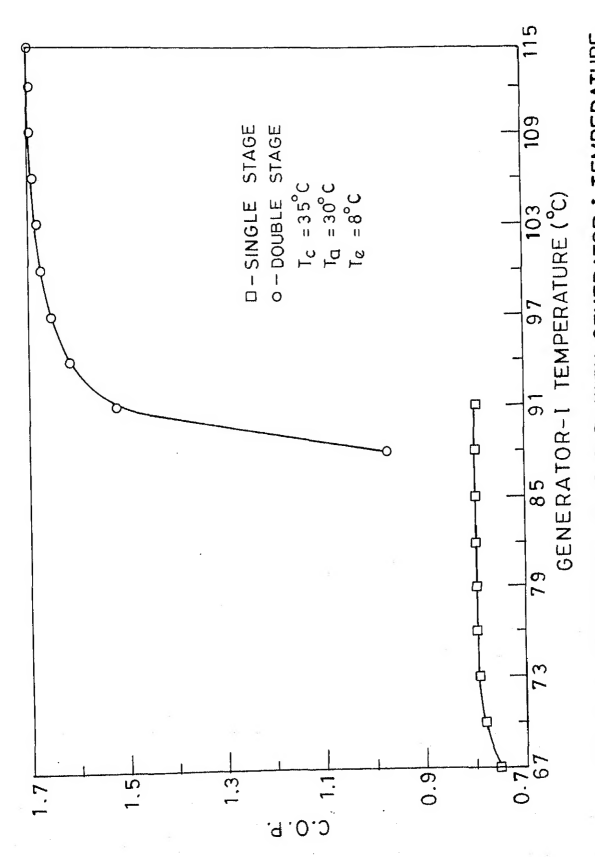


FIG. 5.4 VARIATION OF C.O.P. WITH GENERATOR-I TEMPERATURE.

TABLE -5
Effect of Condenser-Temperature on C.O.P.

	(T _g = 90°	С,	T _a = 25 ^o C ,	T _e = 8 ^o C	$n_1 = n_2 = 0$	0.90)
1	T _c (°C)	3	l _k	5	6	7
1	25.0	60.73	1.758	Cry.	2.732	
2	26.0	60.91	1.753	Cry.	2.580	
3	27.0	61.29	1.745	,	2.445	
4	28.0	61.67	1.736	, ,	2.323	
5	29.0	62,05	1.727	, ,	2.212	
6	30.0	62.26	1.720	, ,	2.111	
7	31.0	62.65	1.709	,,	2.020	
8	32.0	63.05	1.697	Cry .	1.935	
9	33.0	63.28	1.687	.809	1.858	
10.	34.0	63.69	1.671	.807	1.787	•
11	35. 0	64.05	1.658	.805	1.720	

TABLE -6

Effect of Condenser-Temperature on C.O.P.

•	(Tg= 95	oc, T _a =	25°C ,	$T_e = 8^{\circ}C$,	$n_1 = n_2$	= 0.90)
1	T _c (°C)	3	4	5	6	7
1	25.0	63.88	1 . 759	Cry.	3.145	
2	26.0	64.23	1.753	, ,	2.970	·
3	27.0	64.58	1.745	, ,	2.813	
۷Ļ	28.0	64.74	1.740	, ,	2.673	
5	29.0	65.10	1.733	, ,	2,546	
6	<i>5</i> 0.0	65.46	1.725	, ,	2.430	
7	31.0	65.85	1.717	, ,	2.324	
8	<i>3</i> 2.0	66.20	1.708	, , ,	2.227	
9	٥ . وَرَ	66,39	1.702	, ,	2.138	
10	34.0	66.78	1.691	, , ,	2.056	
11	35.0	67,05	1,661	Cry.	1.980	

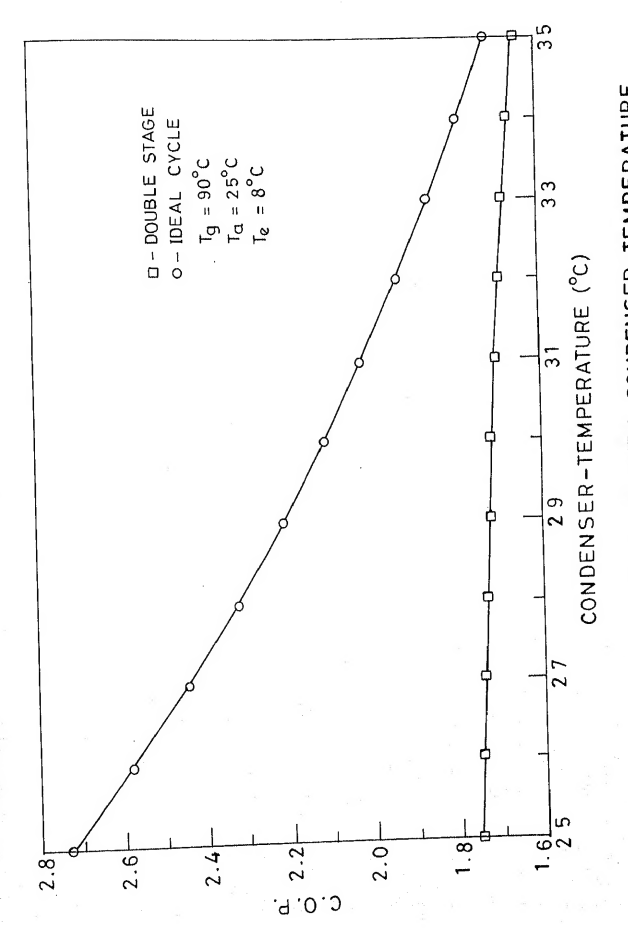


FIG. 5.5 VARIATION OF C.O.P. WITH CONDENSER-TEMPERATURE.

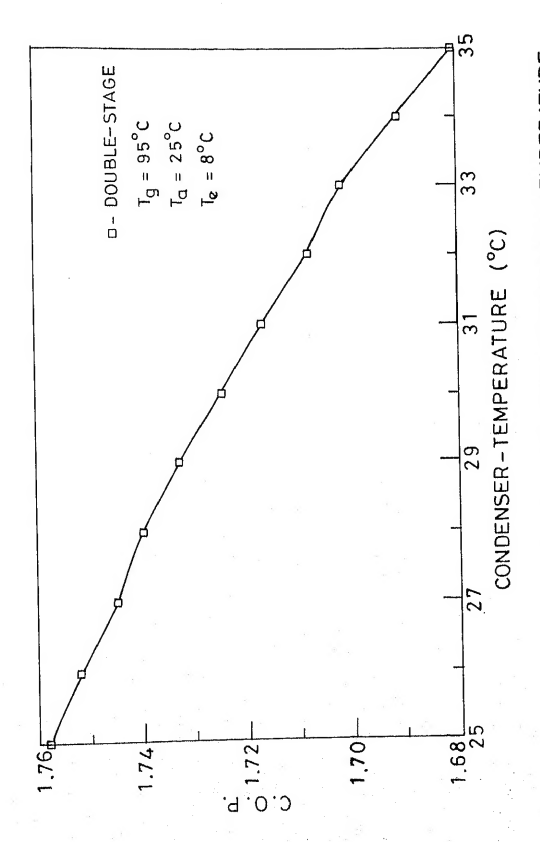


FIG. 5.6 VARIATION OF C.O.P. WITH CONDENSER-TEMPERATURE.

TABLE- 7

Effect of Evaporator-Temperature on C.O.F.

	$(T_g = 85^{\circ}C,$	$T_{c}=$	25°C ,	Ta=20°C	, 7	$n_1 = n_2 = 0.90$
1	T _e (°C)	3.	<i>L</i> _†	5	6	7
1	3.0	57.34	1.728	Cry.	2,278	
2	4.0	57.71	1.743	,,	2.395	
3	5.0	58.26	1.753	5	2.524	
L ₊	6.0	58.83	1.760	, ,	2,666	
5	7.0	59.58	1.765	,,	2.825	
6	8.0	60.16	1.771	, ,	3 . 001	
7	9.0	60,93	1.775	, ,	32.00	
8	10.0	61.23	1.780	,,	3.426	

TABLE- 8

Effect of Evaporator-Temperature on C.O.P.

·····	(T _{g=95} °C	,	$T_{c=25.0C}$,	T _{a=}	20°C ,	$n_1 = n_2 = 0.90$)
1	T _e (°C)	3	4	5	6	7
1	3.0	63.48	1.737	Cry.	2.557	
2	4.0	641 . 09	1.743	,,	2.688	
3	5.0	64.69	1.749	,,	2.833	•
L ₁ .	6.0	65.30	1.754		2.993	
5	7.0	66.12	1.757	2 1	3.171	
6	6.0	66.95	1.760	, ,	3.369	
7	9.0	67.78	1.762	, ,	3.592	•
8	10.0	68.62	1.765	, ,	3.846	

TABLE- 9
Effect of Evaporator-Temperature on C.O.P.

	(T _g = 90°C	,	T _o =30°C ,	$T_a = 2$	25°C, n ₁ =	n ₂ =0.9	90)
1	T _e (°C)	3	4	5	6	7	
1	3 . 0	60.26	1.339	Cry	1.831		
2	4.0	60.62	1,554	,,	1.908		
3.	5.0	60.98	1.635	, ,	1.991		
4	6.0	61.34	1.678	٠,,	2.082		
5	7.0	61.89	1.700	, ,	2.180		
6	8.0	62.26	1.720	,,	2.287		
7	9.0	62.81	1.732	, ,	2,405		-2
8.	10.0	63.37	1.741	, ,	2.534		

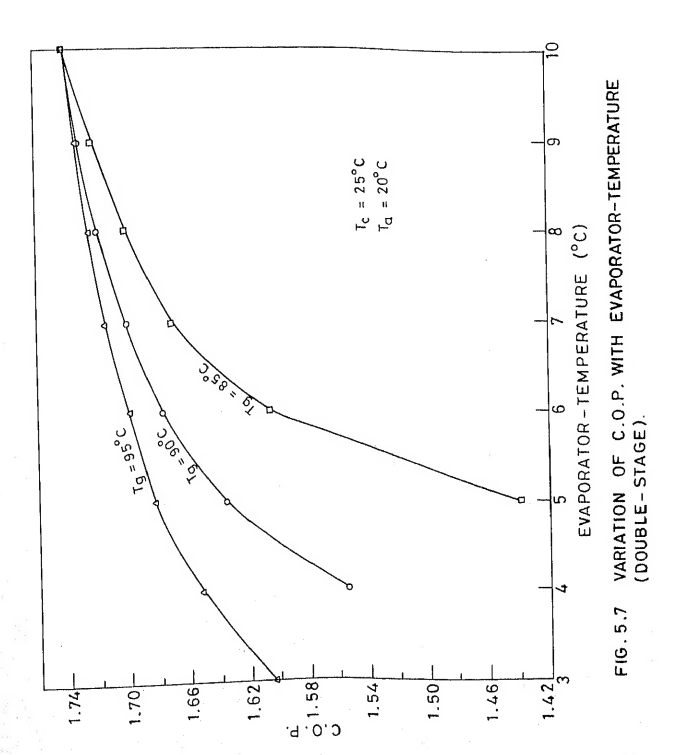


TABLE- 10

Effect of Absorber-Temperature on the C.O.P.

	(T _g =90°C ,	T _c =30°	c, T _e =	=5°C,	$n_1 = n_2 = 0.90$)
1	Ta(°C)	3	4	5	6 7	
1	20.0	63.05	1.719	Cry	2.145	
2	21.0	62,63	1.711	, ,	2.114	
3	22.0	62.07	1.704	,,	2.083	
4	23.0	61.71	1.690	,,	2.052	
5	24.0	61.34	1,669		2.022	
6	25.0	60.98	1.635	,,	1.991	
7	26.0	60.62	1.577	, ,	1.960	
8	27.0	60.26	1.452	, , ,	1.930	
9	28.0	59.91	1.027	,,	1.899	
10	30.0	ONP		.8072	1.869	

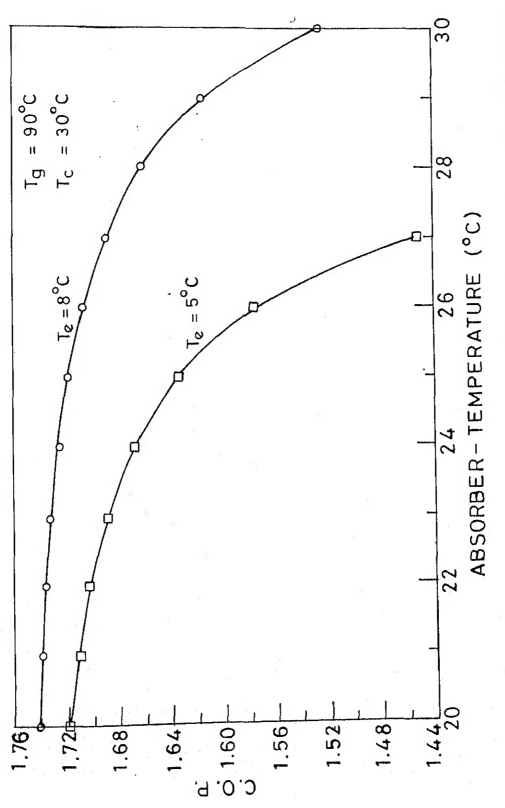


FIG. 5.8 VARIATION OF C.O.P. WITH ABSORBER-TEMPERATURE (DOUBLE STAGE).

TABLE- 11
Effect of Absorber-Temperature on the C.O.F.

	$(T_g = 90^{\circ}C,$	T _c =30°C	, T _e =8 ⁰ C	,	$n_1 = n_2 = 0$.90)
1	T _a (°C)	3	4	5	· 6	7
1	20.0	65.24	1.744	Cry	2.518	
2	21.0	64.88	1.741	,	2.463	
3	22.0	64.31	1.739	, ,	2.428	
L _f	23.0	63.75	1,736	, ,	2.393	
5	24.0	63.18	1.733	,,	2.358	
6	25.0	62.88	1.726	,,	2.322	
7	26.0	62.26	1.720	, ,	2.287	
8	27.0	61.89	1.708	,,	2.252	
9	28.0	61.52	1.690	,,	2.217	
10	29.0	61.16	1.663	, ,	2.182	
11	30.0	60.80	1.524	**	2.111	

TABLE -12

Effect of Heat-Exchanger effectiveness on C.O.P.

		(T _n =90	°C ,	T _c =25°C	$T_a=20$	o°c,	T _e =5°C)		
-	1		ก		3	4					
	1		0.50		61.12	1.325					
	2	¥	0.55		61.12	1.392					
	3	0	0.60		61.315	1.452					
	4		0.65		61.315	1.507					
	5		0.70		61.315	1.556					
	. 6		0.75		61.315	1.598					
	7		0.80		61.509	1.636			* *		
	8		0.85		61.509	1.668		ege.			
	9	,	0.90		61.509	1.698			3 (4) (4) (4)	16	
	10	# B	0.95		61.509	1.726			126		
	11		1.00		61.509	1.788					29.1
						43.4				14	111

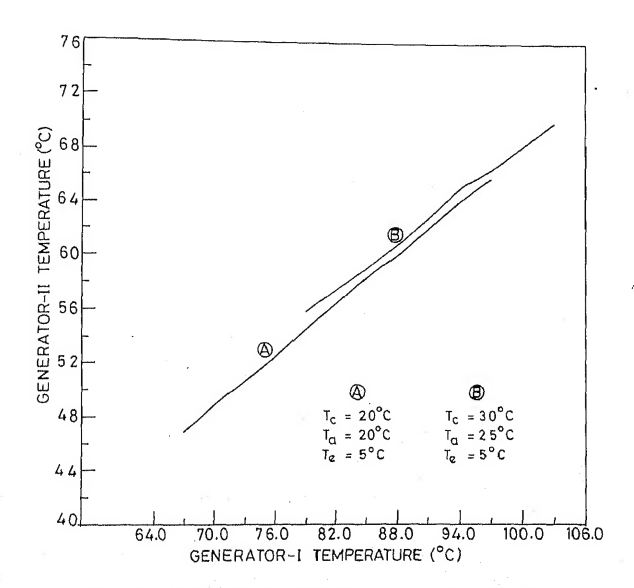


FIG. 5.9 VARIATION OF GENERATOR-II TEMPERATURE WITH GENERATOR-I TEMPERATURE.

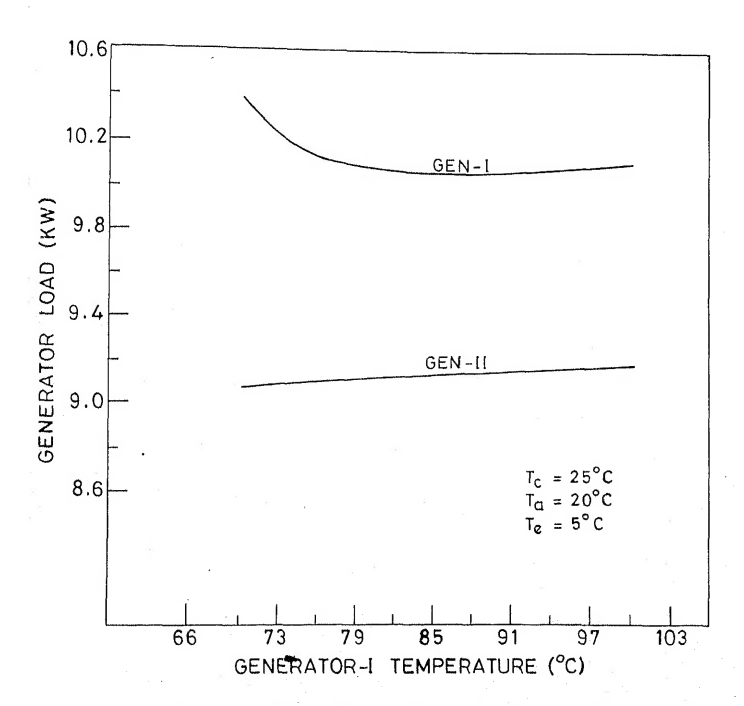


FIG. 5.10 VARIATION OF GENERATORS LOADSWITH GENERA--TOR-I TEMPERATURE.

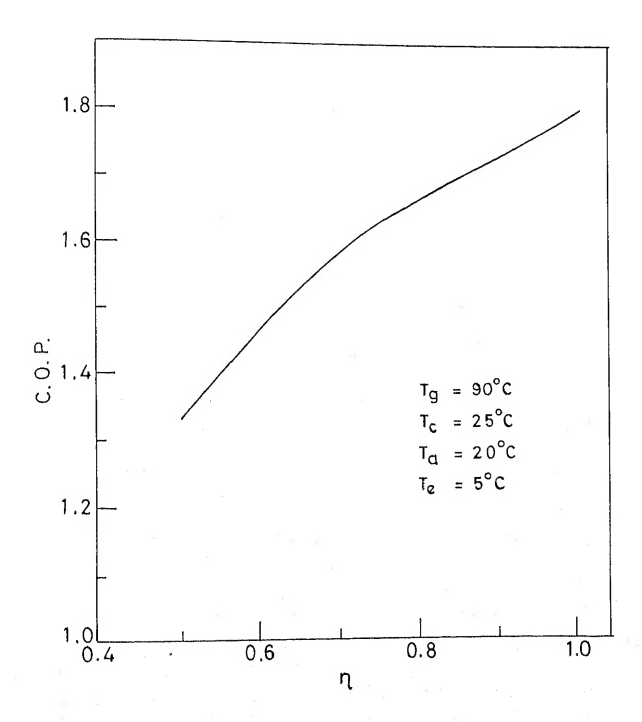


FIG.5.11 VARIATION OF C.O.P. WITH HEAT EXCHAN-GER EFFECTIVENESS.

CONCLUSIONS

An assessment based on steady state thermodynamic analysis of a double effect LiBr-H₂O absorption refrigeration cycle has been made, and its application to space cooling is considered as the main objective. A computer model of the system, based on the simultaneous solution of total mass, material and heat balance equations of different components in the system, has been developed. The proposed FORTRAN programme is very general in nature and simply by changing the input data, the required operational parameters can be determined quite easily.

In view of the results discussed earlier, the following conclusions may be drawn:

(1) The C.O.P. obtained in a double stage-system is quite high (1.785maximum) as compared to single stage (0.84 maximum). However, very high input source-temperature is required for system's operation. This criterion restricts its use to only those places, where very high-temperature steam, bio-gas, hot-water, waste-heat from flue gases and/or advanced solar collectors (evacuated tube), which could give reasonable collector effeciency at high operating

temperatures, are available. However, at locations where only/also high temperature heat sources are available and where the single-stage-system cannot work, a two-stage-system is the solution. Infact, in most of the process industries, the waste heat which is available at quite high temperature, may be successfully utilized for efficient space-cooling purposes.

approaching the ideal value at high condenser temperatures. The practicability of using high condenser-temperatures obviates the necessity of using a cooling tower along with the condenser and an air cooled condenser may be used for the purpose. This helps in a reduction in total cost of the system. Hence, the system would function more effectively in arid regions, where humidity remains low and the ambient temperature is high.

It may, thus, be concluded that a double-stage-vapour-absorption-system is more efficient than a vapour-compression-system in terms of the annual running cost as it does not require compressor for its operation, which consumes most of the energy required for the operation of a vapour-compression-system.

REFERENCES

- 1. Eisenstadt M., Flanijan F.M., Farber F.A.; 'Tests prove feasibility of solar air conditioning '; Heating, piping and Air-Conditioning , vol. 32, no. 11, 1960, pp. 120.
- 2. Verma H.K.; 'Design of one ton solar operated LiBr-H₂O air conditioning system', 'National Solar Energy Convection of Solar Energy Society of India', 'Solar Energy for Rural Development', Bhavnagar, 1978.
- J. Lazzarin R., Rizzan E., Boldrin M.; Scalabrin G.,

 'Performance Prediction of a LiBr-H₂O absorption air
 conditioning utilizing solar energy'; International
 Solar Energy, New Delhi, 1978.
- 4. Alizadeh S., Bahar F., Geoola F.; 'Design and optimization of an absorption refrigeration system operated by Solar Energy', Solar Energy, vol. 22, No. 2, 1979, pp. 149.
- 5. Charters W.W.S., Chen W.D.; 'Some design aspects of air-cooled LiBr-H2O absorption cycle air conditioning system'; SUN-II, vol. 9, 1979, pp. 725.
- 6. Siddiqui A.M., Prasad M., Sahay B.; 'Economic evaluation of bio-gas for optimizing generator temperature in a vapour absorption system'; Energy Conversion and Management, vol. 126, No. 1, 1986, pp. 83.

- 7. Grassie S.L., Sheridan N.R.; 'Modelling of a solar operated absorption air conditioning system with refrigerating storage', Solar Energy, vol. 19, 1973, pp. 691.
- 8. Kaushik S.C., Kumar R.; 'Thermodynamic study of a two stage vapour absorption refrigeration system using NH3 refrigerant with liquid/solid absorbents'; Energy Conversion and Management, vol. 25, No.4, 1985. pp.427.
- 9. Stocker W.F., Jones J.W.; 'Refrigeration and Air Conditioning', McGraw Hill Book Company, 1985.
- 10. Wilbur P.J., Karaki S.; Solar Cooling, The Franklin Institute Press, 1977, pp. 12-13.
- 11. 'Handbook of Air conditioning System Design', Carrier

 Air Conditioning Company, McGraw Hill Book Company.
- 12. ASHRAE Guide and Data Book', Fundamentals and Equipments , 1963.
- 13. Perry, Chilton; 'Chemical Engineering Hand-Book',
 McGraw-Hill Int. Book, 5th edition.
- Duffie J.A.; 'Simulation of a solar heating and cooling system', Solar Energy, vol. 16, 1974, pp. 129.
- 15. Sheridan N.R.; 'Performance of the Brisbane solar house', Solar Energy, vol. 12, 1975, pp. 395.

- 16. Chung R.,; 'A study of a solar air-conditioning';
 Jn of Solar Energy Engg., vol. 85, 1963, pp. 85.
- 17. Colebrook C.F.; Turbulent flow in pipes with particular reference to the transition region between smooth and rough pipe flows; Jn. Inst. Civil Engg., vol. 12, No. 4, 1939, pp. 133-156.
- 18. White F.M.; 'Fluid Mechanics '; McGraw Hill Book Company , 1986, pp. 314.
- 19. 'Standards of the Tubular Exchanger Manufacturers Association', 5th edition, New York, 1970.
- 20. Bogart M.; 'Absorption Refrigeration in Industrial Processes': Houston Gulf Pub. Co., 1981.
- 21. McAdams W.H.; 'Heat Transmission'; McGraw Hill Book Company, 3rd edition, New York, 1974.
- 22. Arora S.C.; 'Refrigeration and Air Conditioning',
 Dhanpat Rai and Sons , 1973.
- 23. Hutchinson F.W.; 'Design of Refrigeration System for Air-Conditioning'; N.Y. Industrial Press, 1969.
- 24. Dossat R.J.; 'Principles of Refrigeration'; New York, Wiley, 1968.
- 25. ASHRAE Handbook of Product Directory', 1977.
- 26. ASHRAE Handbook of Fundamentals , 1981.

APPENDIX -1

State equations for aqueous Lithium-Bromide solution

The well known state equations, relating the solution-temperature and refrigerant-pressure to the refrigerant-temperature, the solution-enthalpy to the solution-temperature, enthalpy of superheated vapour of refrigerant to the generator and condenser-temperatures and the specific heat of the solution to the solution-concentration are given as following \(\int 26.7:\)

$$t_f = A t_f' + B \tag{A-1}$$

$$Log_{10}P = C + \frac{D}{(t'_{f} + 459.72)} + \frac{E}{(t'_{f} + 459.72)^{2}}$$

(A-2)

$$H_{f} = p + q t_{f} + rt_{f}^{2} \tag{A-3}$$

$$H_g = 1.925 T_g - 0.126 T_c + 2500$$
 (A-4)
 $C_{PL} = 4.259 - 0.053843 \times + (2.307E-4) \times 2$ (A-5)

where.

$$A = -2.00755 + 0.16976 \times - (3.133362 E-3) \times^{2} + (1.97668 E-5) \times^{3}$$

$$B = 321.128 -19.322 X + 0.374382 X^{2} - (2.0637 E-3) X^{3}$$

C = 6.21147

 C_{PL} = Specific heat of LiBr-H₂O solution in kJ/kg.K

D = -2886.373

E = -337269.46

 H_f = Enthalpy of solution in Btu/lb

 H_g = Enthalpy of super heated vapour in kJ/kg

 $p = -1015.07 + 79.5387 \times -2.358016 \times^{2} + 0.03031583 \times^{3} - (1.400261 E-4) \times^{4}$

q = 4.68108 - (3.037766E-1) X + (8.44845 E-3) $-(1.047721 E-4) X^{3} + (4.80097 E-7) X^{4}$

r = -4.9107 E-3 + (3.831841 E-4) X-(1.07896 E-5)+ (1.3152 E-7) X³-(5.897 E-10) X⁴

 $t_f = Solution temperature (^{O}F)$

 $40 \le t_f \le 350$ °F

 t'_{f} = Refrigerant temperature (°F) $0 \le t'_{f} \le 230$ °F

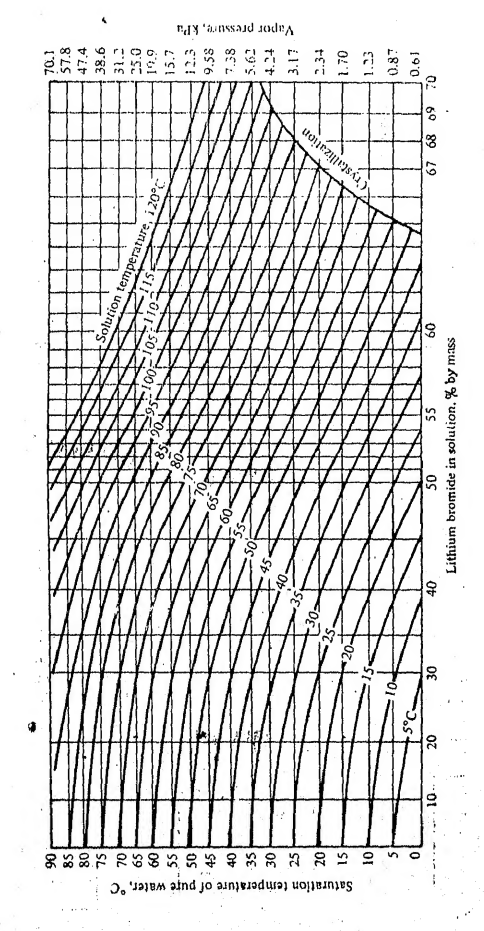
$$= \frac{-2E}{D + \int_{-D^2-4E(C-Log_{10} P)_{-7}^{0.5}}$$

X = Percentage by mass LiBr in LiBr- H_2 0 solution 45 $\% \le X \le 70 \%$

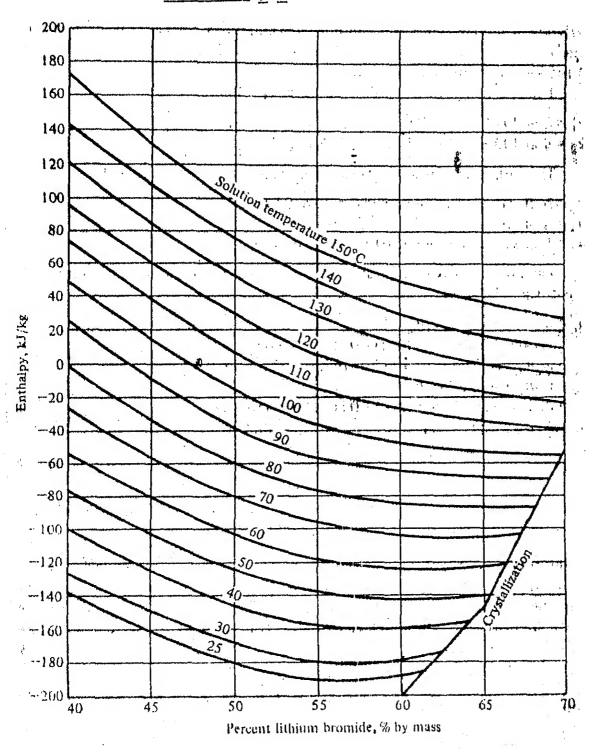
Conversion Factors -

- (1) 1 Btu = 1.0155 kJ
- (2) 1 Ton of Refrigeration = 210 kJ/min
- (3) 1 Psia = 6.9847 k Pa
- (4) 1 Psia = 52.3893 mm of Hg
- (5) 1 Btu/lb= 2.326 kJ/kg

Graph For concentration of LiBr, Vapour Pressure and Saturation Temperature of Pure Water with Solution-Temperature as Variable Parameter /9 7.



Graph for Enthalpy V/S LiBr Concentration with Solution-Temperature oon Variable Parameter /9 /



```
QU2.
004 -
                                                                                                                                                                                                    THE SECTION OF THE SE
                                                                                                                                  THE THE ACTION OF THE STATE LINE WAPPING ADSIRPTION SYSTEM

FIG. 17 ACTION

THE CO. OF THE STATE LINE WAPPING ADSIRPTION SYSTEM

FIND IN THE CO. OF THE SYSTEM IN THE GENER FERH.

PLENT GENERAL TREADERS IN THE GENER FERH.

PLENT GENERAL TREADERS IN THE GENER FERH.

PLENT GENERAL TREADERS IN THE GENERAL THE STATE OF THE GENERATOR

PLENT GENERAL TREADERS IN THE GENERAL THE ACTION OF THE GENERATOR

PLENT GENERAL TREADERS IN THE GENERAL THE GENERATOR

PLENT GENERAL TREADERS IN THE GENERAL THE GENERATOR

PRINT GENERAL TREADERS IN THE GENERAL THE GENERATOR

PRINT GENERAL TREADERS IN THE GENERAL THE EVAPORATOR

PRINT GENERAL TREADERS IN THE GENERAL THE GENERATOR

PRINT GENERAL TREADERS IN THE GENERAL THE GENERAL TREADERS IN 
MO OTH
                                                                                                                                            @1714
OUBLE.
                                                                                                                                             糖いりは
@101 e
0111a
01200
 01 104
 01464
  商生为机构。
270
 # 18C
  1 YOU
  02000
02100
    $220k
    02366
    #2 30E
   0250s
    02700
    CZHE!
    的記り場合
    0.1000
    0316
    03200
      03166
      0350
      0360
      6377
6301
6379
       COUNT
      的名词形
      644622
      网络科
       650
650
       001-
052
               73
         obi.
        (5)
(5)
          C 57* :
                 Same
         (159)
         rice)()
         geit.
          66分
          data de c
           K.G.
           Proc.
           no I · · ·
            er († *).
           6 634
16 7 67
          om i
```

```
is a specification of the Hall
                                             Halifyz, ink
679 L
gazzi i
                                            (5)=0X
67314
                                          % (°) /= / (°)
                                           CHARLEY, SX, CUNCENTRATION OF BUL LEAVING THE ABSURBER= , Fo. 3)
CHARL.
1005 CV:
                                         175=T3(7)
670to
69 /1 m
                                         CALL CONCURS, DIR, OX).
                                         ODBLE.
最も対象
                     10
 SOUCH
$10 144
$11 244
$13 344
                                         ARRESTATION TERMSCRIBER RECIRCULATION PATIO GREGET AND MIEGENERATUR RECIRCULATION RATIO
 2036
                                         AKRS=X(6)/(X(7)-X(6))
                                         $0 /DE
1090
                     87
                                          TO DETERMINE THE HEAT LOADS ON THE VARIOUS COMPUNENTS TANDER ADST MEXT CAUCUDATE THE ENTHAUPTES OF THE FLUIOS TAT VARIOUS STATES.
 111
 $2.200
 27 100
 2111
                                         H(1)=TR(1)*.43+1067.0
H(2)=TRC(2)*4.184/2.326
 11500
11764
                                         11 ( 3 ) = 11 ( 2 ) * 2 ( 3 + 1 ( 6 ) 7 . () ( 6 ) = 1 ( 6 ) = 11 ( 6 ) = 11 ( 6 ) = 11 ( 6 ) = 11 ( 6 ) = 11 ( 6 ) = 11 ( 6 ) = 11 ( 6 ) = 11 ( 6 ) = 11 ( 6 ) = 11 ( 6 ) = 11 ( 6 ) = 11 ( 6 ) = 11 ( 6 ) = 11 ( 6 ) = 11 ( 6 ) = 11 ( 6 ) = 11 ( 6 ) = 11 ( 6 ) = 11 ( 6 ) = 11 ( 6 ) = 11 ( 6 ) = 11 ( 6 ) = 11 ( 6 ) = 11 ( 6 ) = 11 ( 6 ) = 11 ( 6 ) = 11 ( 6 ) = 11 ( 6 ) = 11 ( 6 ) = 11 ( 6 ) = 11 ( 6 ) = 11 ( 6 ) = 11 ( 6 ) = 11 ( 6 ) = 11 ( 6 ) = 11 ( 6 ) = 11 ( 6 ) = 11 ( 6 ) = 11 ( 6 ) = 11 ( 6 ) = 11 ( 6 ) = 11 ( 6 ) = 11 ( 6 ) = 11 ( 6 ) = 11 ( 6 ) = 11 ( 6 ) = 11 ( 6 ) = 11 ( 6 ) = 11 ( 6 ) = 11 ( 6 ) = 11 ( 6 ) = 11 ( 6 ) = 11 ( 6 ) = 11 ( 6 ) = 11 ( 6 ) = 11 ( 6 ) = 11 ( 6 ) = 11 ( 6 ) = 11 ( 6 ) = 11 ( 6 ) = 11 ( 6 ) = 11 ( 6 ) = 11 ( 6 ) = 11 ( 6 ) = 11 ( 6 ) = 11 ( 6 ) = 11 ( 6 ) = 11 ( 6 ) = 11 ( 6 ) = 11 ( 6 ) = 11 ( 6 ) = 11 ( 6 ) = 11 ( 6 ) = 11 ( 6 ) = 11 ( 6 ) = 11 ( 6 ) = 11 ( 6 ) = 11 ( 6 ) = 11 ( 6 ) = 11 ( 6 ) = 11 ( 6 ) = 11 ( 6 ) = 11 ( 6 ) = 11 ( 6 ) = 11 ( 6 ) = 11 ( 6 ) = 11 ( 6 ) = 11 ( 6 ) = 11 ( 6 ) = 11 ( 6 ) = 11 ( 6 ) = 11 ( 6 ) = 11 ( 6 ) = 11 ( 6 ) = 11 ( 6 ) = 11 ( 6 ) = 11 ( 6 ) = 11 ( 6 ) = 11 ( 6 ) = 11 ( 6 ) = 11 ( 6 ) = 11 ( 6 ) = 11 ( 6 ) = 11 ( 6 ) = 11 ( 6 ) = 11 ( 6 ) = 11 ( 6 ) = 11 ( 6 ) = 11 ( 6 ) = 11 ( 6 ) = 11 ( 6 ) = 11 ( 6 ) = 11 ( 6 ) = 11 ( 6 ) = 11 ( 6 ) = 11 ( 6 ) = 11 ( 6 ) = 11 ( 6 ) = 11 ( 6 ) = 11 ( 6 ) = 11 ( 6 ) = 11 ( 6 ) = 11 ( 6 ) = 11 ( 6 ) = 11 ( 6 ) = 11 ( 6 ) = 11 ( 6 ) = 11 ( 6 ) = 11 ( 6 ) = 11 ( 6 ) = 11 ( 6 ) = 11 ( 6 ) = 11 ( 6 ) = 11 ( 6 ) = 11 ( 6 ) = 11 ( 6 ) = 11 ( 6 ) = 11 ( 6 ) = 11 ( 6 ) = 11 ( 6 ) = 11 ( 6 ) = 11 ( 6 ) = 11 ( 6 ) = 11 ( 6 ) = 11 ( 6 ) = 11 ( 6 ) = 11 ( 6 ) = 11 ( 6 ) = 11 ( 6 ) = 11 ( 6 ) = 11 ( 6 ) = 11 ( 6 ) = 11 ( 6 ) = 11 ( 6 ) = 11 ( 6 ) = 11 ( 6 ) = 11 ( 6 ) = 11 ( 6 ) = 11 ( 6 ) = 11 ( 6 ) = 11 ( 6 ) = 11 ( 6 ) = 11 ( 6 ) = 11 ( 6 ) = 11 ( 6 ) = 11 ( 6 ) = 11 ( 6 ) = 11 ( 6 ) = 11 ( 6 ) = 11 ( 6 ) = 11 ( 6 ) = 11 ( 6 ) = 11 ( 6 ) = 11 ( 6 ) = 11 ( 6 ) = 11 ( 6 ) = 11 ( 6 ) = 11 ( 6 ) = 11 ( 6 ) = 11 ( 6 ) = 11
  21804
  1190
                                          ENTHALPY OF THE SOLUTION IS THE FUNCTION OF TEMPERATURE AND
  12000
                                          CONCENTRATION
    2100
                                           $ 2.27
$ 2.34 kg
                                          #(S)=P(X(S))+G(X(S))*TS(S)+R(X(S))*(TS(S)**2)
#(7)=P(X(7))+G(X(7))*TS(7)+R(X(7))*(TS(7)**2)
#(8)=#(7)~ETA1*(H(7)-H(S))
 2240
    2500
                                           吴(日)122(7)
 $201
$27
$201
                      ¢,
                                      DX=X(B)
                                           DH=11(8)
    296
  1334
1334 C
1534 C
    370
    30.
370
     像约(
   S 14 1
     4.4
   471
     3811
     ay.
   gg je
     35
     37
    rën
K
      *6. . .
    to tr
    2 to 15 c
    Lane.
     Telytor
                        4 tu
    in to
    tvis.
    #15.50 )
                                            COP (COP) COP (COP) TO DE PERFORMANCES ', F7.4, Z.5%, 'COP) COP (COP) COP) COP (COP) COP)
      100
```

```
to all Bally DAT
                                                                                                                                                                                                              126 (22, 1)108
                                                                                                                                                                                                (6)=3/
(6)=3/3/
                                                                                                                                                                                                THARTITIES, CONCENTRATION OF BOL LEAVING THE ABSORBERS, F6.3)
                                                                                                                                                                                                    THE TOLY!
                                                                                                                                                                                    CAUL COMOCEDES, DIE, DE)
GELTE (22, 10 ) DE
FORGAT (//, 54, CONCENTRATION OF SOL LEAVING GENERATOR=', F6.3)
                                                                                             10
                                                                                                                                                                                      ARES=37/M1=ABSORBER FECTRODIATION PATIO
GET7=M0/M1=GEDERATOR RECIRCULATION RATIO
                O BURY
   1950
1950
1970
1980
1990
1990
                                                                                                                                                                                    ARROFA(6)/(Xt7)~X(6))
                                                                                                                                                                                      GRE /= ARR5 + 1 .

GRE /= ARR5 + 1 .

GRE /= ARR5 + 1 .

FINAL ELE (22.07) ARR5 = 1 . Fr. 4 . 5 . CRP7 = 1 . Fr. 4 .

FINAL ELE (22.07) ARR5 = 1 . Fr. 4 . 5 . CRP7 = 1 . Fr. 4 . Fr. 
                                                                                              87
                                                                                                                                                                                       TO DETERMINE THE HEAT LUADS ON THE VARIOUS COMPUNENTS THE AUST NEXT CALCULATE THE ENTHALPTES OF THE PLUIDS
        1164
        1200
                  1200
                                                                                                                                                                                         11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
11500
                                                                                                                                                                                      H(1)=TH(1)+.43+1967.9
H(2)=THC(2)+4.184/2.326
                                                                                                                                                                                      11 ( 2 ) - 1 ( 2 ) + 2 ( 3 ) + 3 ( 3 ) + 3 ( 3 ) - 1 ( 3 ) - 1 ( 3 ) - 1 ( 3 ) - 1 ( 3 ) - 1 ( 3 ) - 1 ( 3 ) - 1 ( 3 ) - 1 ( 3 ) - 1 ( 3 ) - 1 ( 3 ) - 1 ( 3 ) - 1 ( 3 ) - 1 ( 3 ) - 1 ( 3 ) - 1 ( 3 ) - 1 ( 3 ) - 1 ( 3 ) - 1 ( 3 ) - 1 ( 3 ) - 1 ( 3 ) - 1 ( 3 ) - 1 ( 3 ) - 1 ( 3 ) - 1 ( 3 ) - 1 ( 3 ) - 1 ( 3 ) - 1 ( 3 ) - 1 ( 3 ) - 1 ( 3 ) - 1 ( 3 ) - 1 ( 3 ) - 1 ( 3 ) - 1 ( 3 ) - 1 ( 3 ) - 1 ( 3 ) - 1 ( 3 ) - 1 ( 3 ) - 1 ( 3 ) - 1 ( 3 ) - 1 ( 3 ) - 1 ( 3 ) - 1 ( 3 ) - 1 ( 3 ) - 1 ( 3 ) - 1 ( 3 ) - 1 ( 3 ) - 1 ( 3 ) - 1 ( 3 ) - 1 ( 3 ) - 1 ( 3 ) - 1 ( 3 ) - 1 ( 3 ) - 1 ( 3 ) - 1 ( 3 ) - 1 ( 3 ) - 1 ( 3 ) - 1 ( 3 ) - 1 ( 3 ) - 1 ( 3 ) - 1 ( 3 ) - 1 ( 3 ) - 1 ( 3 ) - 1 ( 3 ) - 1 ( 3 ) - 1 ( 3 ) - 1 ( 3 ) - 1 ( 3 ) - 1 ( 3 ) - 1 ( 3 ) - 1 ( 3 ) - 1 ( 3 ) - 1 ( 3 ) - 1 ( 3 ) - 1 ( 3 ) - 1 ( 3 ) - 1 ( 3 ) - 1 ( 3 ) - 1 ( 3 ) - 1 ( 3 ) - 1 ( 3 ) - 1 ( 3 ) - 1 ( 3 ) - 1 ( 3 ) - 1 ( 3 ) - 1 ( 3 ) - 1 ( 3 ) - 1 ( 3 ) - 1 ( 3 ) - 1 ( 3 ) - 1 ( 3 ) - 1 ( 3 ) - 1 ( 3 ) - 1 ( 3 ) - 1 ( 3 ) - 1 ( 3 ) - 1 ( 3 ) - 1 ( 3 ) - 1 ( 3 ) - 1 ( 3 ) - 1 ( 3 ) - 1 ( 3 ) - 1 ( 3 ) - 1 ( 3 ) - 1 ( 3 ) - 1 ( 3 ) - 1 ( 3 ) - 1 ( 3 ) - 1 ( 3 ) - 1 ( 3 ) - 1 ( 3 ) - 1 ( 3 ) - 1 ( 3 ) - 1 ( 3 ) - 1 ( 3 ) - 1 ( 3 ) - 1 ( 3 ) - 1 ( 3 ) - 1 ( 3 ) - 1 ( 3 ) - 1 ( 3 ) - 1 ( 3 ) - 1 ( 3 ) - 1 ( 3 ) - 1 ( 3 ) - 1 ( 3 ) - 1 ( 3 ) - 1 ( 3 ) - 1 ( 3 ) - 1 ( 3 ) - 1 ( 3 ) - 1 ( 3 ) - 1 ( 3 ) - 1 ( 3 ) - 1 ( 3 ) - 1 ( 3 ) - 1 ( 3 ) - 1 ( 3 ) - 1 ( 3 ) - 1 ( 3 ) - 1 ( 3 ) - 1 ( 3 ) - 1 ( 3 ) - 1 ( 3 ) - 1 ( 3 ) - 1 ( 3 ) - 1 ( 3 ) - 1 ( 3 ) - 1 ( 3 ) - 1 ( 3 ) - 1 ( 3 ) - 1 ( 3 ) - 1 ( 3 ) - 1 ( 3 ) - 1 ( 3 ) - 1 ( 3 ) - 1 ( 3 ) - 1 ( 3 ) - 1 ( 3 ) - 1 ( 3 ) - 1 ( 3 ) - 1 ( 3 ) - 1 ( 3 ) - 1 ( 3 ) - 1 ( 3 ) - 1 ( 3 ) - 1 ( 3 ) - 1 ( 3 ) - 1 ( 3 ) - 1 ( 3 ) - 1 ( 3 ) - 1 ( 3 ) - 1 ( 3 ) - 1 ( 3 ) - 1 ( 3 ) - 1 ( 3 ) - 1 ( 3 ) - 1 ( 3 ) - 1 ( 3 ) - 1 ( 3 ) - 1 ( 3 ) - 1 ( 3 ) - 1 ( 3 ) - 1 ( 3 ) - 1 ( 3 ) - 1 ( 3 ) - 1 ( 3 ) - 1 ( 3 ) - 1 ( 3 ) - 1 ( 3 ) - 1 ( 3 ) - 1 ( 3 ) - 1 ( 3 ) - 1 ( 3 ) - 1 ( 3 ) - 1 ( 3 ) - 1 ( 3 ) - 1 ( 3 ) - 1 ( 3 ) - 1 ( 3 ) - 1 ( 
         1961
                                                                                                                                                                                         ENTHALPY OF THE SULUTION IS THE FUNCTION OF TEMPERATURE AND
                                                                                                                                                                                         CONCENTRATION
                   2100
                                                                                                                                                                                           2200
                                                                                                                                                                                         担(5)=P(X(5))+Q(X(5))*TS(5)+R(X(5))*(TS(5)**2)
担(7)=P(X(7))+Q(X(7))*TS(7)+R(X(7))*(TS(7)**2)
                   2 3400
                   240
                                                                                                                                                                                         見(数) = ほ(プ) - ETA (*(B(7) - B(5))
                      2500
                                                                                                                                                                                          26.1
                                                                                                                                                                                         DX=X(9)
                      29i (
30i
                                                                                                                                                                                           DHzH(8)
                                                                                                                                                                                       Unit SULTEM(DX,DH,DTS)
TS(W)=DTS
TS(S)=(TS(S)-32.0)*5.0/9.0
$\frac{2}{2}\frac{2}{2}\frac{2}{2}\frac{2}{2}\frac{2}{2}\frac{2}{2}\frac{2}{2}\frac{2}{2}\frac{2}{2}\frac{2}{2}\frac{2}{2}\frac{2}{2}\frac{2}{2}\frac{2}{2}\frac{2}{2}\frac{2}{2}\frac{2}{2}\frac{2}{2}\frac{2}{2}\frac{2}{2}\frac{2}{2}\frac{2}{2}\frac{2}{2}\frac{2}{2}\frac{2}{2}\frac{2}{2}\frac{2}{2}\frac{2}{2}\frac{2}{2}\frac{2}{2}\frac{2}{2}\frac{2}{2}\frac{2}{2}\frac{2}{2}\frac{2}{2}\frac{2}{2}\frac{2}{2}\frac{2}{2}\frac{2}{2}\frac{2}{2}\frac{2}{2}\frac{2}{2}\frac{2}{2}\frac{2}{2}\frac{2}{2}\frac{2}{2}\frac{2}{2}\frac{2}{2}\frac{2}{2}\frac{2}{2}\frac{2}{2}\frac{2}{2}\frac{2}{2}\frac{2}{2}\frac{2}{2}\frac{2}{2}\frac{2}{2}\frac{2}{2}\frac{2}{2}\frac{2}{2}\frac{2}{2}\frac{2}{2}\frac{2}{2}\frac{2}{2}\frac{2}{2}\frac{2}{2}\frac{2}{2}\frac{2}{2}\frac{2}{2}\frac{2}{2}\frac{2}{2}\frac{2}{2}\frac{2}{2}\frac{2}{2}\frac{2}{2}\frac{2}{2}\frac{2}{2}\frac{2}{2}\frac{2}{2}\frac{2}{2}\frac{2}{2}\frac{2}{2}\frac{2}{2}\frac{2}{2}\frac{2}{2}\frac{2}{2}\frac{2}{2}\frac{2}{2}\frac{2}{2}\frac{2}{2}\frac{2}{2}\frac{2}{2}\frac{2}{2}\frac{2}{2}\frac{2}{2}\frac{2}{2}\frac{2}{2}\frac{2}{2}\frac{2}{2}\frac{2}{2}\frac{2}{2}\frac{2}{2}\frac{2}{2}\frac{2}{2}\frac{2}{2}\frac{2}{2}\frac{2}{2}\frac{2}{2}\frac{2}{2}\frac{2}{2}\frac{2}{2}\frac{2}{2}\frac{2}{2}\frac{2}{2}\frac{2}{2}\frac{2}{2}\frac{2}{2}\frac{2}{2}\frac{2}{2}\frac{2}{2}\frac{2}{2}\frac{2}{2}\frac{2}{2}\frac{2}{2}\frac{2}{2}\frac{2}{2}\frac{2}{2}\frac{2}{2}\frac{2}{2}\frac{2}{2}\frac{2}{2}\frac{2}{2}\frac{2}{2}\frac{2}{2}\frac{2}{2}\frac{2}{2}\frac{2}{2}\frac{2}{2}\frac{2}{2}\frac{2}{2}\frac{2}{2}\frac{2}{2}\frac{2}{2}\frac{2}{2}\frac{2}{2}\frac{2}{2}\frac{2}{2}\frac{2}{2}\frac{2}{2}\frac{2}{2}\frac{2}{2}\frac{2}{2}\frac{2}{2}\frac{2}{2}\frac{2}{2}\frac{2}{2}\frac{2}{2}\frac{2}{2}\frac{2}{2}\frac{2}{2}\frac{2}{2}\frac{2}{2}\frac{2}{2}\frac{2}{2}\frac{2}{2}\frac{2}{2}\frac{2}{2}\frac{2}{2}\frac{2}{2}\frac{2}{2}\frac{2}{2}\frac{2}{2}\frac{2}{2}\frac{2}{2}\frac{2}\frac{2}{2}\frac{2}{2}\frac{2}{2}\frac{2}{2}\frac{2}{2}\frac{2}
              1 2 7 . I
                                                                                                                                                                                           別(6) PARR5/GRR7 * (N(7) - H(8))+H(5)
                                                                                                                                                                                           x (6)=X(5)
DX=X(6)
DX=i(6)
            138
                                                                                                                                                                                     DX=X(6)
CAUS SOLTEM(DX,DH,DTS)
TS(0)=TS(6)-32 (0)*5.0(9,0
TS(0)=TS(6)-32 (0)*5.0(9,0
TS(0)=TS(6)-32 (0)*5.0(1) TSC(6)
TSC(0)=TS(6)-32 (0)*5.0(1) TSC(6)
TSC(0)=TS(6)-32 (0)*5.0(1) TSC(6)
TSC(0)=TS(6)-32 (0)*5.0(1) TSC(6)
TSC(0)=TS(0)=TS(0) TSC(0) TSC(0)
TSC(0)=TS(0)=TS(0)=TSC(0)
TSC(0)=TS(0)=TS(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0)=TSC(0
                       12433
                         44
            400
                        1911
1911
                   137
                                                                                                        52
                  1944 I
                   53)
100
                                                                                                                                                                                                 FINE TO AFATER HEATEN, HEATEN, HEATEN, HEATEN HEATEN TO BE KEET AS A THEATEN TO KEET AS A THEAT A SORDER TO KEET AS A THEAT AND THE KEET AND TH
                   7 to $* ..
                   χ (5. (*) ;
                      (1<sup>1</sup>)1 .
                  12
                            Digital.
                                                                                                                                                                                                 Line(1)))

Line(1))

Line(
                                                                                                            100
```

```
ar (Carta) , he ard of the att the
GI, THE
     50
          (1,1,3)
          黄斑龙。
     CCC
美田市 (1)
          REPRODUCTION FOR CANCOLATING CONCENTRATION WHEN TEMPERATURE
          量(红红海)
# Stope St.
     Č
爱想开的。
          SUBSTRUCTATE CUACCUES, DEE, DA)
算13.181.000
          hall stone in
10 9 mg
          TIX SOLVE
     10
          至外的说
EP 2116
77206
12300
DXB=DX-AF/OAF
IF(ABS(DXB-DX).GT.0.01) GO TO 10
29000
          DX=UXII
電門 7節時
17/100
17/100
17/100
          RETURN
          2010g
2010g
          20 JE
20 JE
20 JE
254251
     5
          FECTS**2+B*TS+A-H
          DF=2.0*C*TS+B
DF=2.0*C*TS+B
TSH=TS-F/DF
TF(ABS(TSH-TS).GT.0.01)
TS=TSH
PSETURN
                            10 5
          END:
```

P / H.

```
建打法。
$00 gg a
2013年1
                                                                                           (有点点套套要点标: 新)。然后就想点
簡1951%
$116/2
育り710年
                                                                                                                                            数目的自然
質りりは終
 質えびい
特别其他
1124
1 30%
                                                                                   The control of the standard of the transfer CE,Clus

PRESSED BE TO BE STORM IN THE TREE CE,Clus

PRESSED BE TO THE STORM IN THE TREE CE,Clus

PRESSED BE TO THE STORM IN THE TREE CE,Clus

PRESSED BE TO THE STORM IN THE TREE CE,Clus

FIGURE THE STORM IN THE STORM IN THE TREE CE,Clus

FIGURE THE STORM IN T
 $1500
     1800
1800
     296
     216*
2206
     23
     2 d 0
      25
     2660
  2700
  2.0
2.9
      3000
     350%
       38
      936
    141
        40.2
       301
                                             55
        5F, (
        50
       87.
90
       39 -
        441.74
        (*).
41.
121.
          1 ...
1000
          7 m
          77
```

trop as the absorber pressure

```
C
                                                                                                                                                                                                                                                                                                                                                             Ph(1)2)=28(11)
Ph(1)3/322(12)
                                                                                                                                                                                                                                                                                                                                                                    京安产业现实办法 法宣书节办会 具有法学 朱本泽命不严峻东岛泽海郊城市 电准备不电源表示 医本液溶液 化水准电池 不准不能不安 不管
                                                                                                                                                                                                                                                                                                                                                           Convers the Conve
  0950
0277
                       211
                       1107 = Pa(0)
TATURE OF THE STATE OF THE STAT
                                                                                                                                                                                                                                                                                                                                                           1.4P=05.6
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
112.00
11
                                                                                                                                                                                                                                                                                                                                             Trokes
(c 31 K=1,200
DAX=4(11)
DAX=4(12)
DAX=4
                                                                                                                                                                                                                                                                                                                                                                      ITCH=C
                                                                                                                                                                                                                                                                                                                                                             PH 311 K=1,200
DXX=X(12)
DPRK=PR(1/)
                           15
             25/ 6
27/ 6
27/ 6
22d 6
22/ 6
             2011
                                370
                                39, i
             34
                  .
                       10 (10)
2 (0 (2))
                                         r, 16 i
                                  #Nutr
                           .....
1911
                       en fr
Sister
                    2 44 0 0
2 44 0 0
2 4 0
2 4 0
2 4 0
3 4 0
3 4 0
4 1 0
4 1 0
4 1 0
5 1 0
6 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
7 1 0
                                                                                                                                                                                                                                                                                                                                                                           1(3,2)=4(1+1-4(12))
1(3,4)=(1
1(3,4)=(1,4)
(1,5)=(1,4)=(2)
```

```
(1) (1) (1) (1) (1)
 44).
44)
                    (5)=(1)+(6)
(5)=(-1)+(1)
                  $3 37 E
實用 Te in
         900
進用が行う
                     G SUBROUTINE HAS BEEN USED FOR MATRIX INVERSION ********************************
3 3 m
277 / 100 c
2 (3 th to b)
                  CALL FOGARF (UUU, b, w, 5, y, waspre, 1FAIL)
RECI (1=v(1)
 3950
 71111
                   Cjavtaj
                  (11)=V(3)
(7)=V(3)
(8)=V(5)
((11)=V(6)
                   481=1.0/V(1)
 2700
                   m 1 8 8 3 8 1
20000
20000
 智慧的自
                   TYPE*, ITER, X(11), X(12), X%12

IF((X(12)-XH12), DE.O.15) GO TO 301

F(X(12)-XH12) 301-20
20120
2020
2030
2030
2045
2065
2065
         C
                   303
CC
CC
311
207
208
208
208
208
208
208
                   X(12) = X(12) = 0 al
                   CONTINUE
DCDU=X(7)
                   DENTHEN(7)
CALL SUDTEM (UCON, DENTH, DTEMP)
IS(7)=DTEMP
                   DCOM=X(8)
                   DENTHAM (8)
                   CAGL SOLTÉM (OCOM, DENTH, DTEMP)
                   TS(W)=DTEMP
DCMT=X(11)
DEMTH=H(11)
CALL SOLTEM(OCON, DENTH, DTEMP)
TS(11)=DTEMP
OCON=X(13)
DEMTH=H(13)
CALL SOLTEM(DCON, DENTH, DTEMP)
TS(14)=DTEMP
RMS=10
TMS=FM1*GRR1
2296
                   229
 * '6#
```

```
X(ZZ,5X, TEMP IN DEGR. FORMINETE. ,5X, TEMP. IN DEGR. CENTIC)
                                                                                                 / J=1,5
C(J)=(10(J)~32.J)*5.0/9.Q
U(24,13) J, TM(J), FMC(J)
AT(/,3X,12,7X,FB,J,15x,17.3)
                                                                                   29
                                  53
                                                                                    CTERROR
TPC())=(TR(4)-32.0) *5.0/9.0
JPCTE(24,38) 7.TR(9),TRC(9)
GROAT(Z,3X,12,7X,F8.3,15X,F7.3)
C 53 K=10,13
TSC(X)=(TS(K)-32.0)*5.0/9.0
R1TE(Z4,41) K,TS(K),TSC(K)
GRAT(Z,3X,12,7X,F8.3,15A,F7.3)
                                  38
                                  63
                                                                                            11ADD

16(24,59)

AAT(/,5%, IMPUT TEMPERATURES')

TE(24,64) TSC(10), TEC(3), TSC(6), TPC(4)

TMAT(/,5%, 'SGLUTION TEMP=',F7.3,5%,'CONDENSER TEMP=',F7.3,//,

TMAT(/,5%, 'SGLUTION TEMP=',F7.3,5%,'EVAPORATOR TEMP=',F7.3)

(,'Adsorner Temp=',F7.3,5%,'EVAPORATOR TEMP=',F7.3)
                                  59
                                                                             NLUAD IS THE EVAPORATOR LOAD IN TUNS
REFMAS IS THE REFRIGERANT MASS FLOW RATE THROUGH EVAPORTR
REMAL IS THE REFRIGERANT MASS THROUGH GEN1 KG/MIN
REMAL IS THE REFRIGERANT MASS THROUGH GEN2 IN KG/MIN
REMAL IS THE REFRIGERANT MASS THROUGH GEN2 IN KG/MIN
REMAL IS THE REFRIGERANT MASS THROUGH GEN2 IN KG/MIN
REMAL IS THE REFRIGERANT MASS THROUGH GEN2 IN KG/MIN
                                                                           81
                                   88
                                    93
                                    96
                                     100
                                    150
7.166
                                      130
                                                                                  ና ና ላ ነ ነ ላ ነ ነ ቀ ቀስያት የፊታ የ ነ ነ ነው። ከመደር ነ ነው። ከመደር ነ ነው። ከመደር ነ
                                                                                 CHARLES
CHARLES
CHARLES
```

```
李睿本本本家李本本家华本本本本
                                                                                               PHAT CALCULATING COCENTRATION WHEN
                                                                                                                                                                                                                      REFRIGERANT
                                                                                                                                                    可见的产品以为了的政制 ARE 医内侧的的
海洋中中海原水源水源水源水平水水源水源水源水源水源水源水源水源水源
                                                            李爷水书医哥罗尔科赛马米尔第一张苏格雷米赛马车等海岸华斯群
 31,000
                                                            BURELLE CONCERTS, DIR, UA)
 3011
                                                        1. 14 = 4 3 . 13
                                                         ប្រាំ≖ស់%់
                            10
                                                        DAMPA AL SILIAM Z. OC/55*018-DTS)*((0.16976*DTR)*19.322)*DX*
1(0.1374382*(U18*3.133362k-3))*(UX**2)*(UTR*1.97668E-5*2.0637
11-3)*(UX**3)
(AF=((0.16976*DTR)*49.322)*Z.0*(0.374382*(DTR*3.133362E-3))*DX
1+3.4*(DTR*1.97668E-5*Z.0037E*3)*(DX**2)
UX: =0X*AF/UX*
IF(ABS(OX*-OX)/UX*.GT.0.00001) GO TU 10
 3031 1
 30300
 3555
 3556.
 3 8 Mar
 3081 .
 37000
                                                        DX=UXD
3736
3736
3736
3756
3756
3776
3776
3776
                                                        BETURN
                                                         Ç
                                                         SUBSUUTINE FOR CONCENTRATION WHEN SULUTION TEAPERATURE AND
                                                         以1999年1998年,在1998年,1998年,1999年,1999年,1998年,1998年,1998年,1998年,1998年,1998年,1998年,1998年,1998年,1998年,1998年,1998年,1
1998年,1988年,1998年,1998年,1998年,1998年,1998年,1998年,1998年,1998年,1998年,1998年,1998年,1998年,1998年,1998年,1998年,1998年,1
                                                         SHERDUTINE GISECULPTS, DPR, OTF, DX)
                                                        379
  3000
 3010
                                                        DTR=-2.0*E/(D+SQRT(D**2-((4.0*E)*(C-DPRES))))-459.72
  jv2...
  3031
                                                         DX=UXN
  30 4
                             15
                                                        DX=UAN

AFF=(321.128-2.00/55*DTR-DTS)+(0.16976*DTR)-19.322)*DX

1+(0.3/4382-(DTR*3.133362E-3))*(DX**2)+(DTR*1.97668E-5-2.0637

1E-3)*(DX**3)

DAFF=((0.16976*DTR)-19.322)+2.0*(0.374382-(DTR*3.133362E-3))*DX

1+3.0*(DTR*1.97668E-5-2.0637E-3)*(DX**2)

DXN=DX*AFF/DAFF
  38568
  30600
  38716
  3000
 3901
                                                         TR (ABS(DXN-OX), GT.0.001) GO TO 15
     9100
  3 9 2 ( )
3 9 3 ( )
                                                        SUBROUTINE WHEN PRESSURE AND CONCENTRATION ARE KNUITEMPERATURE IS TO CALCULATE WHICH EMABLES US TO FIRMESSURE OF GENERATURE!
 KNOWA SOLUTION
TO FIND HIGH
                                                        SUBSTITUTE HIGHPRODX DPRH OTRE 2FT OTS APPRIL 1.97668E-5)

**-1.00755+0.16976*DXA-(3.133362F-3)*(DXX**2)+(1.97668E-5)

**(OXX**3)

B=321.128-19.322*DXX+0.374382*(DXX**2)-(2.0637E-3)*(DXX**3)

OPRESS ADD TO (DPRR)

C=0.25147

D=-2030.371

E=-11776*.46

OTS APPIRED TO (D**2)-((4.0*E)*(C*DPRESS))))-469.72

D**-2030.371

C=-11776*.46

OTS C=A*DIR + B+6.49

OTS C=A*DIR + B+6.4
  405
  4000
  410
 Til, Piliti
  427
   427
   4...
   431.
  4 7 1
4 7 1
4 7 1
4 7
4 7
4 7
```

ľ.

APPENDIX - 6

Condenser Data for Vapour Compression System (For Refrigerant-Freon 22), taken from the Central Air Conditioning Unit, I.I.T. Kanpur.

(2) **ater Pressure	(1)	Refrigerant Pressure	230 psig
Outlet 85°F (4) Pipe size of condenser ID=5/8 dia(16mm) OD= 18 mm Length=10,126 tubes (5) Pipe material (6) No. of working hours 24	(2)	water Pressure	
OD= 18 mm Length=10,126 tubes (5) Pipe material Cu tubes, steel shell (6) No. of working hours 24 (7) No. of condensing units 9 (8) Compressors 7 Nos90 h.p. 2 Nos100 h.p. (9) Condenser Pumps 3 Nos 30 h.p. 1 No 25 h.p. (10) Full Load 720 tons	(3)	Water temperature	
(6) No. of working hours 24 (7) No. of condensing units 9 (8) Compressors 7 Nos90 h.p. 2 Nos100 h.p. (9) Condenser Pumps 3 Nos 30 h.p. 1 No 25 h.p. (10) Full Load 720 tons	(4)	Pipe size of condenser	OD= 18 mm
(7) No. of condensing units (8) Compressors 7 Nos90 h.p. 2 Nos100 h.p. (9) Condenser Pumps 3 Nos 30 h.p. 1 No 25 h.p. (10) Full Load 720 tons	(5)	Pipe material	Cu tubes, steel shell
(8) Compressors 7 Nos90 h.p. 2 Nos100 h.p. (9) Condenser Pumps 3 Nos 30 h.p. 1 No 25 h.p. (10) Full Load 720 tons	(6)	No. of working hours	24
2 Nos100 h.p. (9) Condenser Pumps 3 Nos 30 h.p. 1 No 25 h.p. (10) Full Load 720 tons	(7)	No. of condensing units	9
(9) Condenser Pumps 3 Nos 30 h.p. 1 No 25 h.p. (10) Full Load 720 tons	(8)	Compressors	7 Nos90 h.p.
1 No 25 h.p. (10) Full Load 720 tons			2 Nos100 h.p.
	(9)	Condenser Pumps	
(11) Operating Load (Month-May) 550 tons.	(10)	Full Load	720 tons
	(11)	Operating Load (Month-May)	550 tons.

Material (New)	ε in mm
Revetted Steel	0.9-9.0
Concrete	0.3-3.0
Wood Stave	0.18-0.9
Cast Iron	0.26
Galvanized Iron	0.15
Commercial Steel	0.046
Drawn Tubing	0.0015

```
m 11 5/21 5 (1)
                                                                                                                                                                                                                                                                                           A. CHRESHAR SCIENCE & PHOTO MECHANICS)
1.197
 10.41.3
                                                                                                                                                                                                                                                                                                                                                                                                                                                                              UMPARING
AND TOO
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   OPERATIONAL COST OF STAGE VAPOUR ABSORPTION SYS
                                                                                                                                                                              ** * 1 . . .
 MAL.
 # 1.3:
$15°
10:5
· 11/1/
 1411
 720:1
  123 .
                                                                                                                                                                                                 9201
 @25t
 626/1
  $271.
 簡望在665
                                                                                                                                                                                    TRARESERS TO STARE TO STARE THE STARE STAR
  029
 Q 3000.
 $ 31 m
                                                                                                                                                                                    PRESI=23.445
PRESI=0.483
PRESI=0.483
ARRI=25.5
ARRI=110/M1
 03200
 otto
             34
 自35的标
                                                                                                                                                                                   REF 1-110(-1.0

REF 12-16EN2M*GBR2

POSE1-1REF 12+43.d05)*(PRE52-PRES1)/(1.65*1000.0*.80*60.0)
             346
                                                                                                                                                                                 POWER1=3.0*POWP1
COST1=PUWR1*24.0*0.05
COST1 IS THE COST ASSOCIATED TO RATSE THE PRESSURE LEVEL OBLY
OTHER CUSTS INCLUDES PRESSURE DROP DUE TO FRICTIONAL LOSSES
 039
             OUR
ζ,
                                                                                                                                                                             OTHER COSTS INCLUDES PRESSURE DROP DUE TO FRICTIONAL WARREST CONTROL OF THE COSTS IN CHOOSEN TO GIV ALLOWABLE SOLUTION VELOCITY 2M/SEC CALL FRIC (EPIS, DIA, FRFACT)
FI=FRFACT
DELPI=FI*15.0*2.0*2.0*1650.0*1000./(2.0*2.5*25.4)
PRESSURE LOSSES DUE TO BEND
GE=0.45
VEL=2.6
DELPI=VEL**2*1650.0*GF/2.0
DELPI=VEL**2*1650.0*GF/2.0
DELPI=O.013
DIA=1.0*25.4
CALL PRIC (EPIS, DIA, FRFACT)
FI=FRFACT
DELPI=FI2*3.0*1.5*1.5*1650.0*1009.0/(2.0*25.4)
             ¥7
                                                                                           (
             50
           19534
  OU
                                                                                                                                                                                   PUMINISTOR PI+DELP2+DELV3)*CREFM12+43.87/C1000.0*1650.0*.80*60.03)
PUMER REPONDE *1.0
CUST 1: PUMER REPONDE *1.0
CUST PUMER REPONDE *1.0
CUST
   602
     t till:
                                                                                                                                                                                The contract of the contract o
    f . 11 je e
   137 · ·
   MG (7' .
   r ny: i
   6 J.W.
  ECCHERON, DOLA, EXELUTION OF A CONTROL OF A
```

```
=| | | | | | |
                                                                                                                                             STATE TO THE TENED TO THE TOTAL TO THE TOTAL TOTAL TOTAL TOTAL TOTAL TO THE TOTAL TO
     0.7914
  100mm
                                                                                                                                              21.244
21.344
      豊村養利
                                                                                                                                             Pere /=0cb87*4A16Pa/(990.0*100g.0)
PCVE87=PAGE7*2.0
CLS17=PAGE7*0.65*2.0
TYCE*, COST: COST
     145
     16700
       10900
     10900
     11000
   212km
213km
214km
                                                                                                                                               VICERPICED AD TROOD.

JAN 185 - FEBRUARY COLL

JAN 185 - FEBRUARY TO TECHT, YECGZ, DAISA, SAVING

TO BOAT (10A, YUNG) LOEB. COST VAP. COMP SYSTEM=', F8.1,//, 10X,

FY VEHROY UPER. COST FOR VAP. ABS. SYST=', F8.1,//, 24X, DAILY

250 VILG EU ES=', F8.2,//, 24x, YEARDY SAVING IN RS=', F8.1)
     11500
    11000
   117/1/
1180//
1180//
120///
                                                                                                                                               ស្តីទីក្រុង
ស្ត្រីក្រុ
                                                                                                                                              SUBROUTINE FUR FINDING OUT FRICTIONAL FACTOR WITH SOLUTION SUBROUTINE FUR (EPIS, DIA, FREACT)
 ASDIG/EPIS
                                                                                                                                           DEK=1050.0
VEL=2.0
VIS=1.454*0.01*0.1
PEYNO=DEN*YEL*DIAZVIS
B=REXROZA
22220122
22220122
12220122
12221
                                                                                                                                          IF CABSTERNEW FREACT) . LE. O. OOGGOO1) GO TO 10
                                                                                                                                          IF CARS(FRHEN-FRFACT).LE.O.OOGOO1) GO TO 10

FRFACT=FRHEN

FRFACT=FRHEN

HETURA

FRICTORA

SUBROUTINE FOR FRICTOR FACTOR WHEN BATER TO FLOWING

SUBROUTINE WAFRICIOERIS, ROTA, FRIER)
 36
200
200
200
200
201
201
201
201
                                                                      10
                                                                                                                                           DEN=1000.0
YIS=0.080773
VEL=1.5
A=001A/DEPI8
REYNU=DEN#WEU*DOLAZVIS
 2 A ()
  1044
144
                                                                                                                                         REYNDADER*VEL*DOLAXVIS
SERVANIA
FIFASOUR (FRIER)+1.0/(1.14+2.0*ALOG10(A)-2.0*ALOG10(1.0+9.3)
FIRETS-SURFERNIAL)

DESCRIPTION (FRIER)///
DESCRIPTION (FRIERA)+((1.0/(1.14+2.0*ALOG10(A)-2.4*ALOG
(1.0+0.3/(2.4*CINKIEA))))*(2.0/(1.0+9.3/(6*SONT(FRIERA)))
FRIERASURT (FRIERASURT (FRIERA)))
FRIERASURT (FRIERASURT (FRIERASURT))
FRIERASURT (FRIERASURT)
FRIERASURT (FRIERASUR
   250
          57. .
   195.
   150
    2570
           561
    9 3 JA
                                                                      5, ,,
```

CHE-1907-M-MEH-COM